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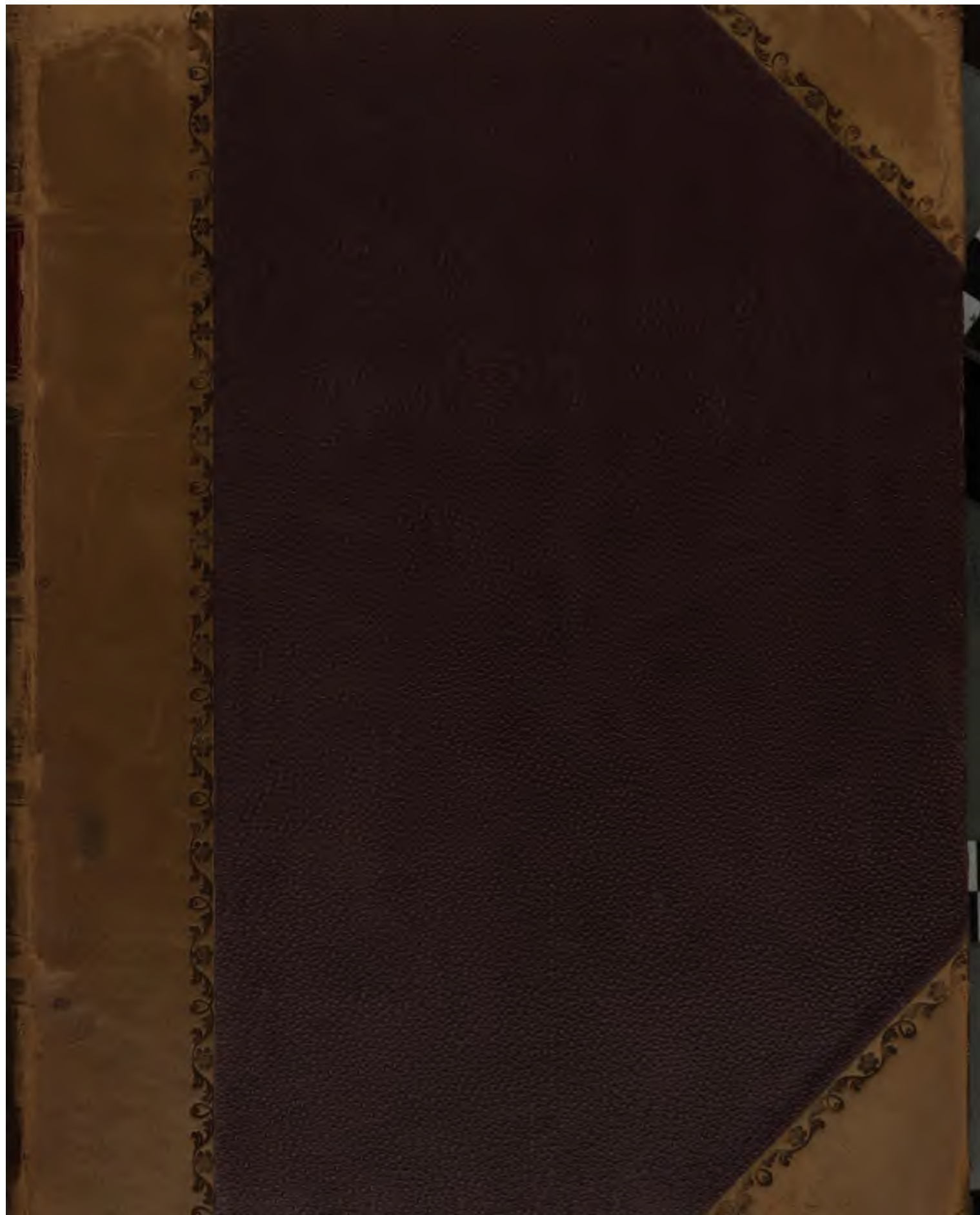
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IN ALL ITS APPLICATIONS TO
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EDITED BY
WILLIAM CROOKES, F.R.S., &c.

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THE CHEMICAL NEWS.

VOLUME XXXIX.

EDITED BY WILLIAM CROOKES, F.R.S., &c.

No. 997.—JANUARY 3, 1879.

DISCUSSION OF THE WORKING HYPOTHESIS THAT THE SO-CALLED ELEMENTS ARE COMPOUND BODIES.*

By J. NORMAN LOCKYER, F.R.S.

It is known to many Fellows of the Society that I have for the last four years been engaged upon the preparation of a map of the solar spectrum on a large scale, the work including a comparison of the Fraunhofer lines with those visible in the spectrum of the vapour of each of the metallic elements in the electric arc.

To give an idea of the thoroughness of the work, at all events in intention, I may state that the complete spectrum of the sun, on the scale of the working map, will be half a furlong long; that to map the metallic lines and purify the spectra in the manner which has already been described to the Society, more than 100,000 observations have been made and about two thousand photographs taken.

In some of these photographs we have vapours compared with the sun; in others vapours compared with each other; and others again have been taken to show which lines are long and which are short in the spectra.

I may state, in way of reminder, that the process of purification consisted in this:—When, for instance, an impurity of manganese was searched for in iron, if the longest line of Mn was absent, the short lines must also be absent on the hypothesis that the elements are elementary; if the longest line were present, then the impurity was traced down to the shortest line present.

The Hypothesis that the Elements are Simple Bodies does not include all the Phenomena.

The final reduction of the photographs of all the metallic elements in the region 39-40—a reduction I began in the early part of the present year, and which has taken six months, summarised all the observations of metallic spectra compared with the Fraunhofer lines accumulated during the whole period of observation. Now this reduction has shown me that the hypothesis that identical lines in different spectra are due to impurities is not sufficient. I shall show in detail in a subsequent paper the hopeless confusion in which I have been landed. I limit myself on the present occasion to giving tables showing how the hypothesis deals with the spectra of iron and titanium.

We find short line coincidences between many metals, the impurities of which have been eliminated or in which the freedom from mutual impurity has been demonstrated by the absence of the longest lines.

Evidences of Celestial Dissociation.

It is five years since I first pointed out that there are many facts and many trains of thought suggested by solar and stellar physics which point to another hypothesis—namely, *that the elements themselves, or at all events some of them, are compound bodies.*

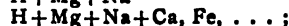
In a letter written to M. Dumas, December 3, 1873, and printed in the *Comptes Rendus*, I thus summarised a memoir which has since appeared in the *Philosophical Transactions*.

“Il semble que plus une étoile est chaude plus son spectre est simple, et que les éléments métalliques se font voir dans l'ordre de leurs poids atomiques.”

“Ainsi nous avons :

“1. Des étoiles très-brillantes où nous ne voyons que l'hydrogène, *en quantité énorme*, et le magnésium ;

“Des étoiles plus froides, comme notre Soleil, où nous trouvons :—



dans ces étoiles, pas de métalloïdes ;

“Des étoiles plus froides encore, dans lesquelles tous les éléments métalliques sont associés, où leurs lignes ne sont plus visibles, et où nous n'avons que les spectres des métalloïdes et des composés.

“4. Plus une étoile est âgée, plus l'hydrogène libre disparaît ; sur la terre, nous ne trouvons plus d'hydrogène en liberté.

“Il me semble que ces faits sont les preuves de plusieurs idées émises par vous. J'ai pensé que nous pouvions imaginer une ‘dissociation céleste,’ qui continue le travail de nos fourneaux, et que les métalloïdes sont des composés qui sont dissociés par la température solaire, pendant que les éléments métalliques monatomiques, dont les poids atomiques sont les moindres, sont précisément ceux qui résistent, même à la température des étoiles les plus chaudes.”

Before I proceed further, I should state that while observations of the sun have since shown that calcium should be introduced between hydrogen and magnesium for that luminary, Dr. Huggins's photographs have demon-

* Paper read at the Royal Society, December 12, 1878.

* This referred to the old numbers in which $Mg = 12$. $Na = 23$.

{ CHEMICAL NEWS,
{ January 3, 1879.

[illegible]

TABLE II.—FINAL REDUCTION—TITANIUM.

Intensity in Sun.	Wave-length and length of line.	Coincidences with Short Lines.									
	39										
	0.000	Zr									
1	3	4									
	0.048	Th									
4	3	4									
	10.10		Mn	Ce	Li						
5	5		4	5	3						
	13.60					Va					
2	3					4					
	19.15				Ce						
5	8				4						
	20.50						U		I.a		
4	3						3		3		
	23.68					Va					
3	2					3					
	37.13	Th		Ce							
3	5	4		4							
	47.75						Fe				
2	1						2				
	57.22	Zr							Rh		
2	1	1							3		
	61.75						U				
4	2						3				
	63.35				Li					Ta	
3	2				3					5	
	80.83						Fe				
2	1						2				
	81.52									Mo	
3	2									3	
	89.22		Mn							Cr	
1	1		4							4	
	97.98					Va					
2	longest					longest					

strated the same fact for the stars, so that in the present state of our knowledge, independent of all hypotheses, the facts may be represented as follows, the symbol indicating the spectrum in which the lines are visible.

Hottest stars	Fluted	H + Ca + Mg	
Sun	Bands of	H + Ca + Mg + Na + Fe	
Cooler Stars	Lines of	— — — Mg + Na + Fe + Bi + Hg	
		— — — — — — — — — —	
Coollest..		— — — — — — — — — —	Metalloids

Following out these views, I some time since communicated a paper to the Society on the spectrum of calcium, to which I shall refer more expressly in the sequel.

Differentiation of the Phenomena to be Observed on the Two Hypotheses.

When the reductions of the observations made on metallic spectra, on the hypothesis that the elements were really elementary, had landed me in the state of utter confusion to which I have already referred, I at once made up my mind to try the other hypothesis, and therefore at once sought for a critical differentiation of the phenomena on the two hypotheses.

Obviously the first thing to be done was to inquire whether one hypothesis would explain these short line coincidences which remained after the reduction of all the observations on the other. Calling for simplicity sake the short lines common to many spectra basic lines,

the new hypothesis, to be of any value, should present us with a state of things in which basic molecules representing bases of the so-called elements should give us their lines, varying in intensity from one condition to another, the conditions representing various compoundings.

Suppose A to contain B as an impurity and as an element, what will be the difference in the spectroscopic result?

A in both cases will have a spectrum of its own;

B as an impurity will add its lines according to the amount of impurity, as I have shown in previous papers.

B as an element will add its lines according to the amount of dissociation, as I have also shown.

The difference in the phenomena, therefore, will be that, with gradually increasing temperature, the spectrum of A will fade, if it be a compound body, as it will be increasingly dissociated, and it will not fade if it be a simple one.

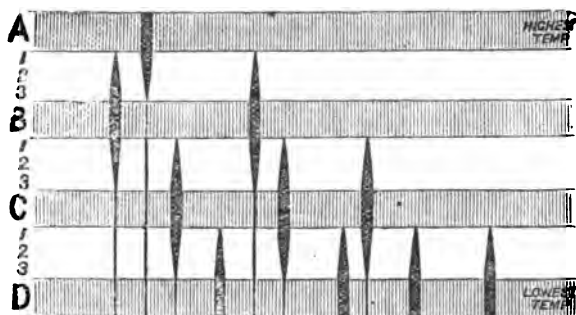
Again, on the hypothesis that A is a compound body, that is, one compounded of at least two similar or dissimilar molecular groupings, then the longest lines at one temperature will not be the longest at another, the whole fabric of "impurity elimination," based upon the assumed single molecular grouping, falls to pieces, and the origin of the basic lines is at once evident.

This may be rendered clearer by some general considerations of another order.

General Considerations.

Let us assume a series of furnaces A . . . D, of which A is the hottest.

FIG. 1.*



Let us further assume that in A there exists a substance α by itself competent to form a compound body β by union with itself or with something else when the temperature is lowered.

Then we may imagine a furnace B in which this compound body exists alone. The spectrum of the compound β would be the only one visible in B, as the spectrum of the assumed elementary body α would be the only one visible in A.

A lower temperature furnace C will provide us with a more compound substance γ , and the same considerations will hold good.

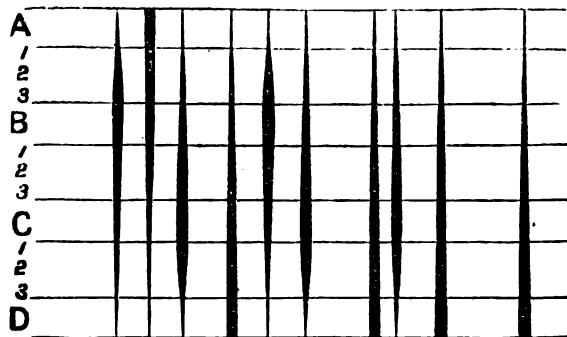
Now if into the furnace A we throw some of this doubly compounded body γ we shall get at first an integration of the three spectra to which I have drawn attention; the lines of γ will first be thickest, then those of β , and finally α would exist alone, and the spectrum would be reduced to one of the utmost simplicity.

This is not the only conclusion to be drawn from these considerations. Although we have by hypothesis β , γ , and δ all higher, that is, more compound forms of α , and although the strong lines in the diagram may represent the true spectra of these substances in the furnaces B, C, and D respectively, yet, in consequence of incomplete dissociation, the strong lines of β will be seen in furnace C, and the strong lines of γ will be seen in furnace D, *all as thin lines*. Thus, although in C we have no line which is not represented in D, the intensities of the lines in C and D are entirely changed.

In short, the line of α strong in A is *basic* in B, C, and D, the lines of β strong in B are *basic* in C and D, and so on.

I have prepared another diagram which represents the facts on the supposition that the furnace A, instead of having a temperature sufficient to dissociate β , γ , and δ into α is far below that stage, although higher than B.

FIG. 2.



* The figures between the hypothetical spectra point to the gradual change as the spectrum is observed near the temperature of each of the furnaces.

It will be seen from this diagram that then the only difference in the spectra of the bodies existing in the four furnaces would consist merely in the relative thicknesses of the lines. The spectrum of the substances as they exist in A would contain as many lines as would the spectrum of the substances as they exist in D; each line would in turn be *basic* in the whole series of furnaces instead of in one or two only.

Application of these General Considerations to Impurity Elimination.

Now let us suppose that in the last diagram (FIG. 2) the four furnaces represent the spectra of say, iron, broken up into different finenesses by successive stages of heat. It is first of all abundantly clear that the relative thicknesses of the iron lines observed will vary according as the temperature resembles that of A, B, C, or D. The positions in the spectra will be the same, but the intensities will vary; this is the point. *The longest lines, represented in the diagram by the thickest ones, will vary as we pass from one temperature to another.* It is on this ground that I have before stated that the whole fabric of impurity elimination must fall to pieces on such an hypothesis. Let us suppose, for instance, that manganese is a compound of the form of iron represented in furnace B, with something else; and suppose again that the photograph of iron which I compare with manganese represents the spectrum of the vapour at the temperature of the furnace D. To eliminate the impurity of iron in manganese, as I have eliminated it, we begin the search by looking for the longest and strongest lines shown in the photograph of iron, in the photograph of manganese taken under the same conditions. I do not find these lines. I say, therefore, that there is no impurity of iron in manganese, but although the longest iron lines are not there, some of the fainter basic ones are. This I hold to be the explanation of the apparent confusion in which we are landed on the supposition that the elements are elementary.

Application of these Considerations to Known Compounds.

Now to apply this reasoning to the dissociation of a known compound body into its elements—

A compound body, such as a salt of calcium, has as definite a spectrum as a simple one; but while the spectrum of the metal itself consists of lines, the number and thickness of some of which increase with increased quantity, the spectrum of the compound consists in the main of channelled spaces and bands, which increase in like manner.

In short, the molecules of a simple body and a compound one are affected in the same manner by quantity in so far as their spectra are concerned; *in other words, both spectra have their long and short lines*, the lines in the spectrum of the element being represented by bands or fluted lines in the spectrum of the compound; and in each case the greatest simplicity of the spectrum depends upon the smallest quantity, and the greatest complexity (a continuous spectrum) upon the greatest.

The heat required to act upon such a compound as a salt of calcium so as to render its spectrum visible, dissociates the compound according to its volatility; the number of true metallic lines which thus appear is a measure of the quantity of the metal resulting from the dissociation, and as the metal lines increase in number, the compound bands thin out.

I have shown in previous papers how we have been led to the conclusion that binary compounds have spectra of their own, and how this idea has been established by considerations having for a basis the observations of the long and short lines.

It is absolutely similar observations and similar reasoning which I have to bring forward in discussing the compound nature of the chemical elements themselves.

In a paper communicated to the Royal Society in 1874,

referring, among other matters, to the reversal of some lines in the solar spectrum, I remarked "—

"It is obvious that greater attention will have to be given to the precise character as well as to the position of each of the Fraunhofer lines, in the thickness of which I have already observed several anomalies. I may refer more particularly at present to the two H lines 3933 and 3968 belonging to calcium, which are much thicker in all photographs of the solar spectrum [I might have added that they were by far the thickest lines in the solar spectrum] than the largest calcium line of this region (4226'3), this latter being invariably thicker than the H lines in all photographs of the calcium spectrum, and remaining, moreover, visible in the spectrum of substances containing calcium in such small quantities as not to show any traces of the H lines.

"How far this and similar variations between photographic records and the solar spectrum are due to causes incident to the photographic record itself, or to variations in the intensities of the various molecular vibrations under solar and terrestrial conditions, are questions which up to the present time I have been unable to discuss."

An Objection Discussed.

I was careful at the very commencement of this paper to point out that the conclusions I have advanced are based upon the analogies furnished by those bodies which, by common consent and beyond cavil and discussion, are compound bodies. Indeed, had I not been careful to urge this point the remark might have been made that the various changes in the spectra to which I shall draw attention are not the results of successive dissociations, but are effects due to putting the same mass into different kinds of vibration or of producing the vibration in different ways. Thus the many high notes, both true and false, which can be produced out of a bell with or without its fundamental one, might have been put forward as analogous with those spectral lines which are produced at different degrees of temperature with or without the line, due to each substance when vibrating visibly with the lowest temperature. To this argument, however, if it were brought forward, the reply would be that it proves too much. If it demonstrates that the *h* hydrogen line in the sun is produced by the same molecular grouping of hydrogen as that which gives us two green lines only when the weakest possible spark is taken in hydrogen inclosed in a large glass globe, it also proves that calcium is identical with its salts. For we can get the spectrum of any of the salts alone without its common base, calcium, as we can get the green lines of hydrogen without the red one.

I submit, therefore, that the argument founded on the overtones of a sounding body, such as a bell, cannot be urged by any one who believes in the existence of any compound bodies at all, because there is no spectroscopic break between acknowledged compounds and the supposed elementary bodies. The spectroscopic differences between calcium itself at different temperatures is, as I shall show, as great as when we pass from known compounds of calcium to calcium itself. There is a perfect continuity of phenomena from one end of the scale of temperature to the other.

Inquiry into the Probable Arrangement of the Basic Molecules.

As the results obtained from the above considerations seemed to be so far satisfactory, inasmuch as they at once furnished an explanation of the *basic lines* actually observed, the inquiry seemed worthy of being carried to a further stage.

The next point I considered was to obtain a clear mental view of the manner in which, on the principle of evolution, various bases might now be formed, and then become basic themselves.

It did not seem unnatural that the bases should increase their complexity by a process of continual multiplication, the factor being 1, 2, or even 3, if conditions were available under which the temperature of their environment should decrease, as we imagined it to do from the furnace A down to furnace D. This would bring about a condition of molecular complexity in which the proportion of the molecular weight of a substance so produced in a combination with another substance would go on continually increasing.

Another method of increasing molecular complexity would be represented by the addition of molecules of different origins. Representing the first method by $A + A$, we could represent the second by $A + B$. A variation of the last process would consist in a still further complexity being brought about by the addition of another molecule of B, so that instead of $(A + B)_2$ merely, we should have $A + B_2$.

Of these three processes the first one seemed that which it was possible to attack under the best conditions, because the consideration of impurities was eliminated; the prior work has left no doubt upon the mind about such and such lines being due to calcium, others to iron, and so forth. That is to say, they are visible in the spectra of these substances as a rule. The inquiry took this form. Granting that these lines are special to such and such a substance, does each become basic in turn as the temperature is changed?

I therefore began the search by reviewing the evidence concerning calcium and seeing if hydrogen, iron, and lithium behaved in the same way.

(To be continued)

ANALYSIS OF BOILER FEED-WATERS.

By W. F. K. STOCK, F.C.S., F.I.C.

BEING frequently engaged in the analysis of boiler feed-waters, I have found, as a result of several years' experience, that unless the person to whom a report is submitted happens to be possessed of considerable chemical knowledge, a statement giving an exhaustive analysis of a water residue is apt to be very much more puzzling than edifying, and, as a matter of fact, such an analysis is by no means necessary to the purpose of selecting waters for boiler use. A much more simple method of procedure has stood one in good stead in several difficult cases, and is the one I always follow, of course with modifications to meet special requirements.

Most of the readers of the CHEMICAL NEWS will be familiar with the characteristics of a good boiler water, but for the sake of making more complete the description of the mode of working, I may be allowed to point them out as follow:—

1. Freedom from any very appreciable quantity of suspended mineral matter.
2. Absence of any trace of mineral acids or of acid salts, or corrosive salts of any kind.
3. Absence of oily or fatty substances.
4. A good boiler water should not contain more than 30 grains solids per gallon, and not more than half of this should precipitate on boiling under pressure.

Some little consideration is here due to the statements thus made. For example, the amount of suspended mineral matter a boiler water may contain is, to a great extent, governed by the quantities of carbonates of lime and magnesia, and sulphate of lime it contains, and by the manner in which the boiler is fed. If a water gave a coherent deposit on boiling, I should feel bound to object to more than 2 or 3 grains of mineral matters per gallon in suspension, as tending to augment and harden such deposit; and, again, if a boiler were to be fed without subsidence or filiation, the same objection would hold

good on account of damage to pumping and feed apparatus. With respect to mineral acids or corrosive salts my opinion is that any such contamination ought to condemn a water utterly for boiler purposes, unless simple and effective means could be adopted for their neutralisation.

The action of oils and fats has of late received considerable attention in connection with boiler waters. For some time the opinions respecting their influence were extremely various, but it is now pretty generally acknowledged that their presence is of no possible good, and may lead to very serious harm. The following case, which came under my own observation in March, 1876, may be of interest in this connection. A boiler at a large ironworks in the Cleveland district was being fed with Tees water (a most excellent boiler water), which was slightly heated on its way to the boiler, by the exhaust steam from the blowing engines, with which steam, however, it never came into actual contact. It was observed that after the boiler had been at work for about two years, the feed pipe supplying it with the warmed Tees water became leaky in places not far removed from a flanged joint, the perforations from which the water oozed being like very small pin-holes. The boiler was neighbour to another, having a cast-iron feed pipe, in which no leak whatever existed. The faulty pipe was removed, and put into my hands for examination. I first of all paid attention to the feed water, and, finding no clue there, turned my attentions upon the pipe itself. I had some difficulty in detecting the exterior perforations, but where they occurred the interior of the pipe was eaten away to such an extent that a walnut would have laid in the holes. These holes were filled with spongy masses of what appeared to be ferric oxide, but which upon analysis gave the following results:—

Ferric oxide	=	66.91	per cent
Ferrous	=	23.69	"
Aluminic	=	trace	"
Calcic	=	0.60	"
Magnesian	=	0.70	"
Sulphur trioxide	=	0.22	"
Phosphoric pentoxide	=	0.24	"
Water	=	1.60	"
Organic matter	=	5.31	"
Insoluble	=	1.30	"

100.57

The organic matter yielded oleic acid in quantity, on treatment with ether, and this, coupled with the presence of both ferric and ferrous oxides, and the comparative ease with which wrought-iron filings are attacked by oleic acid in presence of water, when heated under pressure, led me to adopt the following explanation of the corrosion. The presence of the fatty matter must, under the circumstances, be attributed to accident; most probably this had occurred during erection of plant. The first effect of the action of the fatty acid upon the metal was the production of a ferrous salt, which was oxidised by the oxygen existing in solution in the water, the iron only being raised in the scale of oxidation, the fat acid being liberated, and seizing upon another portion of metal, which was again oxidised, and so the action became local and continuous. I do not believe the cast metal pipe would have suffered at all under such circumstances, because the graphite it contained would have protected it from corrosion, and under my advice cast pipes were supplied to all the boilers as a preventive measure. It is seldom, indeed, or never that all the features I have pointed out would occur in any individual sample of water, and as a rule the most the analyst has to do is to indicate what amount of scale-forming matter the sample contains; and when it is called to mind what terrible accidents have occurred, and are constantly happening, through the unadvised use of hard waters, it becomes a matter of the utmost importance to have some reliable and simple method of making this

point clear to steam users. I have placed the limit figure in a water analysis for boiler purposes at 15 per gallon, which number is the result of observation, partly, and of inquiry amongst practical men using which were known to me. Any greater contents lead to more frequent blowing off and chipping than is the and loss of time is the consequence. The evil becomes much worse if attention is withheld, and the ultimate result is local overheating, unequal expansion and traction, culminating either in sudden destruction of boiler, with its too-often attendant horrors, or the boiler becomes utterly leaky and unmanageable. The following is the method I have devised and adopted as embodying the whole of the foregoing, and as furnishing in short time most reliable information as to the character of any given water generally:—

1. The suspended matter is determined by filtering 700 cc. of the water through tared papers. The residue is washed, dried at 110°C., and weighed, burnt and weighed again. The weight in grammes gives grains per gallon, and the difference between first and second weighings gives the suspended matter.
2. The examination for free mineral acids consists in testing the water, which must be rendered clear by filtration, if necessary, with dilute coccolite tincture, and determining the acid so found by decinormal soda. If corrosive salts are suspected they are best arrived at by the evaporative method. A large volume of the water (700 to 1500 c.c.) is concentrated by the action of the concentrated liquid, a weighed strip of pure iron, which should be previously washed with boiled distilled water, is then with strong alcohol, dried, and weighed. The nature of the corrosive substance is, of course, given in the report, with any other information the analyst can supply regarding it.
3. Oily and fatty matters occasion a milky appearance in water containing them, which disappears on treatment with a moderate quantity of ether. The amount of ether or fat is determined by evaporating 350 c.c. of the water-bath (with the addition of one or two drops of dilute sulphuric acid) to about 70 c.c. The residual acid liquid is cooled, digested with ether in a stoppered tube about two centimetres diameter. The ethereal portion is decanted into a watch glass, the acid liquid washed with more ether, which is added to that in the dish, and the etheral solutions evaporated over the water-bath and weighed; and this weight, reported along with the weight of caustic soda needed to render the water clear, or oil, found harmless. It is only in condensed waters that such substances are found as a rule, but however, grease should occur in water, giving lime and magnesia salts, sufficient soda would be required for their removal also.
4. The proportions of solids deposited on boiling under pressure and solids retaining solubility on boiling under pressure, are found by taking an observation of the total solid matter the water contains, by evaporation of 70 c.c. in platinum, and dry at 100°C., is quite sufficiently accurate. It is necessary to boil 700 c.c. of the water for 24 hours in a flask connected by an india-rubber stopper, with an inverted Liebig's condenser fed with cold water. The boiled water is then cooled to temperature at which it was measured, the condenser has received proper attention, and appreciable diminution of volume will have taken place. The cooled water is run through a dry filter, and 70 c.c. are evaporated at 100°C. as before. The difference between the weights of the residues gives solids deposited on simply boiling. We have now to add to this the calcium su-

contained in the boiled and filtered water, because calcium sulphate is practically insoluble at a pressure of one atmosphere of steam. The calcic oxide is determined in 350 c.c. of the boiled and filtered water, calculated into dry sulphate, and added on to solids deposited on boiling. The difference between the number so obtained and the total solids gives solids retaining solubility on boiling under pressure, which solids are only of account in as far as they augment the boiling-point of the residual water. It will of course be obvious that if a water contains free acids, except carbonic, that latter part of the process will be omitted.

I have always viewed the boiling of a water in a thin glass flask as a very excellent method of judging of its scale-forming qualities, and it is quite remarkable to note various peculiarities of deposits so obtained.

In reporting an analysis of boiler water made by this method I adopt the following tabulation:—

	Grains per gal.	Pounds per 100 galls.
Solids deposited by boiling under pressure ..	—	—
Solids remaining in solution on boiling under pressure	—	—

Then follow remarks as to the analyst's opinion of the water generally. I need hardly say that it is the bounden duty of every one who attempts the analysis of boiler water to make himself well acquainted with the construction and manner of working the different makes of boilers in ordinary use, and also to study well the reactions of the various salts natural waters contain when subject to heat and pressure combined. When one views the evidence produced before coroners' juries dealing with cases of death from boiler explosions, it is often very painfully apparent that there is a great deal yet to be done by chemists in the tracing of effect to cause, the information afforded being often of the most meagre quality, and a field of inquiry embracing the salvation of life and property is surely one in which honour is not wanting.

Laboratory and Assay Office, Darlington,
December 19, 1878.

ON THE METHOD OF EXTRACTING GOLD, SILVER, AND OTHER METALS FROM PYRITES.*

By W. A. DIXON, F.C.S., Cor. Mem. Nat. Hist. Soc. Glasgow.
(Concluded from p. 303).

THE method of treatment of the concentrated ore, or regulus, is the same whether the sulphides are rich in the precious metals or not, but requires variation according to—First, the presence or absence of copper; second, the proportion of copper; and, third, the presence or absence of lead.

To begin with the simplest case, viz., with pyrites free from copper. The apparatus required consists of—1, a roaster; 2, a reverberatory furnace; 3, an arsenic flue, (if the pyrites are arsenical); 4, a leaden combination chamber; 5, a leaden condensing tower; 6, a series of lixiviating tanks and coolers. It is best to construct the furnace, roaster, arsenic flue, converting chamber, and tower in one line, so that the waste heat from the reverberatory furnace heats the sole of the roaster and converting chamber. The reverberatory furnace is constructed in the usual manner, but the sole is made simply of brick, and flat, with an opening in the centre or side, through which the charge may be raked out into an iron hopper

waggon. The roaster is built as a muffle, with a sole of brick or cast-iron plates laid at a slight incline, to facilitate the transference of the charges. At the lower end there is a depression of about 6 inches, forming a recess, which extends half way over the reverberatory furnace, and has an opening which can be closed with a slide, through which the contents of the recess may be at once transferred to the furnace. At the end of the roaster is an arsenic flue if required. Farther on is the combination chamber, built of sheet-lead, supported by an external framing of wood; the sole is of sheet-lead, supported on iron plates over the flue. The sole is divided into eight stages, each of which is an inch higher than the one nearer the roaster. Beyond the combination chamber is a condensing tower, which may be built of brick, laid in a putty of clay and coal-tar, or, better, sheet-lead, supported by an external framing of wood. The condenser is filled with coke or pebbles, supported on iron bars covered with lead, allowing a free space for the entry of the gases evolved in the roaster. Surmounting the coke is a perforated sheet of lead, with suitable openings for the escape of the residual gases, which may be conducted by a pipe to a chimney. On the perforated lead plate water is delivered at intervals by a self-acting tumbler, kept supplied by a cistern. The water escaping through the holes in the lead plate is uniformly distributed over the coke, and trickles downwards, escaping over the sole of the combination chamber, flowing over each stage in succession, and from the lowest to a wooden cooler, whence it is again pumped to the cistern.

The roaster should have two or three small fireplaces at intervals underneath the sole, to get it to a working heat, which may be closed when this is attained, the flame from the reverberatory furnace, mixed with a sufficient quantity of air through openings in the flue, then supplying sufficient heat.

The roaster sole being heated to a dull red heat, 2 cwts. of the dressed pyrites, or ground regulus, is mixed with from 10 to 15 per cent of previously roasted ore, and charged into the upper end of the roaster, so that it occupies about 2 feet of the length, and spreads across the sole, which it should cover to the depth of one and a half inches. The ore soon becomes ignited, evolving sulphurous, sulphuric, and arsenious oxides. The last is condensed in the arsenic flue, and the two former pass through the combination chamber to the condensing tower, and are there absorbed by the descending water.

In an hour's time the charge is moved 2 feet to the left, and a second charge of the mixture is placed in the space cleared. At the end of another hour the first charge is moved 2 feet to the left, the second to the space cleared, and a third is introduced, and so on, until at the end of twenty-four hours the sole of the roaster is covered. The upper part of the roaster should be at a very dull red heat, whilst the lower should be sufficiently hot to decompose any sulphate of iron formed.

On the sole of the recess from 35 to 40 lbs. of coal-dust, charcoal-dust, or other carbonaceous matter is now spread, and the calcined residue from the first charge is turned over on top of it, each charge in the roaster is moved downwards, and a fresh charge of mixture introduced. In another hour a similar quantity of carbon is spread on top of the charge in the recess, and the second charge is turned over on top, and so on until eight successive charges of roasted residue and carbon are in the recess.

The contents of the recess are then transferred to the reverberatory furnace through the opening; the whole is well stirred up, and spread over the sole. The furnace is closed, and kept at a moderate red-heat for eight hours, the furnace being kept full of a smoky flame to assist reduction. The oxides are thus reduced to the metallic state, and the heat should be kept so low as to prevent the reduced iron from agglutinating into masses. At the end of eight hours the reduced metal is withdrawn from the furnace at the opening into the hopper waggon, which is immediately closed, so as to prevent access of air. The

* Read before the Royal Society of N.S.W. August 1, 1877.

furnace is again charged from the recess, which has meanwhile been filled.

When the hopper is cool enough to be handled, the contents are rapidly transferred to a vessel containing water, best by placing the waggon over the vessel and withdrawing a slide in its bottom, so as at once to thoroughly wet and cool the contents.

One-eighth part, or thereabouts, of the cooled metal is now placed in the upper stage of the combination chamber, over which water from the condenser, charged with sulphurous and sulphuric acids, is flowing. In one hour the charge is moved to the second stage, and a second charge is introduced, and so on, until in eight hours the chamber is filled, and that furnace charge exhausted. The metal is rapidly acted on by the acids, and converted into sulphate, sulphite, and hyposulphite of iron; but by the combined action of the air and water, assisted by the heat from the flue, the first largely predominates. The charge on the lowest stage is now removed, and may be at once lixiviated with water; but it is best to keep it for a few days in a moist state, to finish the conversion into sulphate. It is then lixiviated with water, which removes sulphate of iron (also zinc, nickel, and cobalt if present), and leaves a residue containing the gold and silver, mixed with quartz, excess of carbon, and free sulphur. The carbon and sulphur are then burned off, and the gold and silver separated from the quartz by washing, amalgamating, or otherwise. If the extraction of the sulphate of iron is effected with boiling water, and the liquor run into coolers, that salt may be obtained in a marketable condition; or, the crystals being calcined at a dull-red heat, yield a fine red oxide of iron suitable for painting, and sulphuric acid, which may be condensed.

If the ores contain copper in small proportion only, the roasting, reduction, and solution of the iron are conducted precisely as above described, using, however, as little carbon as possible for the reduction, so as to obtain the residue nearly free therefrom. The copper is then found in the residue principally as sulphide. This residue is roasted at a dull red heat, and the copper is extracted by treatment with condenser liquor and crystallisation, as described below, the gold and silver being obtained from the residue as before.

Ores or sulphides containing much copper are roasted at a dull red heat, the mixing with roasted ore, roasting, and condensation of the gases being conducted as before. The well-roasted ore is withdrawn from the furnace and cooled. The cooled residue is then shaken into a lixiviating tank partly filled with a solution of sulphate of copper, containing sulphuric acid and sulphurous acid, obtained as below described; the mixture is allowed to digest, and treated with successive portions of the solid solution until the escaping liquor contains free sulphuric acid. The lixiviation is then carried on with a cold saturated solution of sulphate of copper from the coolers, which is made boiling hot in a leaden or copper boiler, until the specific gravity of the entering and escaping solutions is the same. The whole of the copper liquors are run into wooden coolers, where crystals of the sulphates of copper and iron are deposited. When the lixiviation has been carried as far as possible with the copper solution, it is drawn from the tank until it stands only an inch or so above the solid contents, and 12 inches of water are carefully floated on top, and the drawing off the copper solution continued from below until the water is only an inch above the solid contents, when a second wash is run on, and in the same manner a third if necessary. The copper liquors are run to the coolers as long as they mark above 20° of Twaddell, below that strength they are run to a separate tank to be used for the first wash of another lot.

If the ore contains silver, a little is found in solution in the copper liquor, and is separated therefrom by filtering it through a bed or beds of cement copper, or, better, of precipitated sulphide of copper, before running the liquor to the coolers. The silver is recovered from time to time

by toasting the precipitate, extracting the copper by condenser liquor, and melting the residual silver.

The residue in the lixiviating tanks is drained, dried, mixed with one-fourth of its weight of carbon, reduced, and otherwise treated as above described, to obtain the gold and remaining silver.

The crystals of sulphates of copper and iron in the coolers are removed from time to time, drained and dried. One ton of the dried crystals is charged into the muffle furnace, and there exposed to a fully cherry-red heat, so as to convert the whole of the sulphates into oxides. The sulphurous and sulphuric acids evolved are conveyed to the condensing tower, which is supplied with sulphate of copper solution from the coolers, slightly diluted with the weaker wash liquor by which the acids are condensed, and used for extracting roasted ore. When vapours are no longer evolved the calcined residue is removed from the muffle. A similar charge of dried sulphate is heated in the muffle furnace to a dull red, so as to convert the sulphate of iron into oxide, and when fumes are no longer evolved the charge is raked out, mixed with $\frac{1}{2}$ cwts. of coal or charcoal dust, and charged into the reverberatory furnace, where it is melted, when sulphurous and other gases are evolved, causing the mass to boil. When boiling ceases the oxides are added to the charge, and the melting heat continued until the whole is in a tranquil fusion, when the slags are raked off and the rough copper run into moulds.

If the ore contains lead it is found in the residue containing the gold and silver; and if present in sufficient quantity the residue may be smelted, and the gold and silver recovered by cupellation.

If not present in sufficient quantity to smelt, but still so much as to interfere with the amalgamation, the residue is roasted, treated with a little condenser liquor, washed, and the lead extracted with solution of caustic soda, when the gold and silver may be amalgamated. The solution of lead in caustic soda is mixed with sawdust or carbon, evaporated to dryness, heated strongly, and the carbonate of soda dissolved out with water, and again rendered caustic by lime, when the lead remains as an insoluble residue mixed with carbon.

The advantages of this mode of treatment are that the sulphides are entirely got rid of, whilst if through inattention in the roasting some sulphides remain, only the small proportion that has escaped requires to be re-roasted, instead of the whole mass of ore as is usually the case. In the extraction of copper from the sulphides the whole of the copper may be obtained in the form of crystals, from which the copper may be recovered without evaporation of liquors; and in the whole of the operations, with the exception of smelting for copper, the temperatures are so low that the cheapest materials may be used for the construction of furnaces. The temperature being low, the loss of silver by volatilisation is reduced to a minimum, whilst there is absolutely no loss of gold, except through the careless spilling of the material. Lastly, as neither salt, iron, or other material is consumed in the process, a large source of expense in all previous wet methods of treatment is avoided; and the process is adapted for use wherever the ores are found.

PROCEEDINGS OF SOCIETIES.

MANCHESTER LITERARY AND PHILOSOPHICAL SOCIETY.

Ordinary Meeting, November 26, 1878.

J. P. JOULE, D.C.L., LL.D., F.R.S., &c., President, in the Chair.

"On some Improved Methods of Producing and Regulating Electric Light," Part II., by HENRY WILDE.

In a former communication to the Society I directed attention to the fact that when the electric light is produced from the ends of two carbon pencils placed parallel to each other, if the strength of the electric current, the thickness of the carbons, and the distance between them are rightly proportioned, the carbons will burn steadily downwards until they are wholly consumed, without any insulating material between them. To initiate the light by this method it is necessary to complete the electric circuit between the carbons by means of some conducting substance, which volatilises on the passage of the current, and establishes the electric arc between the points.

When a number of such lights are produced simultaneously from the same source of electricity, any interruption in the continuity of the current extinguishes all the lights in the same circuit, and each pair of carbons requires to be re-primed before the lights can again be established. This defect, as will be obvious, would cause great inconvenience when the lights are not easily accessible, or are at considerable distances apart.

In the course of my experiments it was observed that when the electric circuit was completed at the bottom of a pair of carbons close to the holders, the arc immediately ascended to the points, where it remained so long as the current was transmitted. My first impression of this peculiar action of the arc was, that it was due to the ascending current of hot air by which it was surrounded. This, however, was found not to be the cause, as the arc travelled towards the points in whatever position the carbons were placed, whether horizontally or vertically in an inverted position. Moreover, when a pair of carbons were held in the middle of the holders, the arc travelled upwards or downwards towards the points, according as the circuit was established above or below the holders. The action was in fact recognised to be the same as that which determines the propagation of an electric current through two rectilinear and parallel conductors submerged in contact with the terrestrial bed, which was described by me in the *Philosophical Magazine*, August, 1868.

In all the arrangements in general use for regulating the electric light, the carbon pencils are placed in the same straight line, and end to end. When the light is required, the ends are brought into momentary contact, and are then separated a short distance to enable the arc to form between them. The peculiar behaviour of the electric arc when the carbons are placed parallel to each other suggested to me the means of lighting the carbons automatically, notwithstanding the fact that they could only be made to approach each other by a motion laterally, and to come into contact at their adjacent sides. To accomplish this object one of the carbon holders is articulated or hinged to a small base plate of cast-iron, which is so constructed as to become an electro-magnet when coiled with a few turns of insulated wire. The carbon holder is made in the form of a right-angled lever, to the short horizontal limb of which is fixed an armature placed over the poles of the electro-magnet. When the movable and fixed carbon holders are brought into juxtaposition, and the carbons inserted in them, the upper parts of the two carbons are always in contact when no current is transmitted through them.

The contact between the carbons is maintained by means of an antagonistic spring inserted in a recess in one of the poles of the electro-magnet, and reaching on the under side of the armature. One extremity of the coil of the electro-magnet is in metallic connection with the base of the carbon holder, while the other extremity of the coil is in connection with the terminal screw at the base of the instrument, from which it is insulated. The coils of the electro-magnet are thus placed in the same circuit as the carbon pencils.

When the alternating current from an electro-magnetic induction machine is transmitted through the carbons, the electro-magnet attracts the armature and separates the upper ends of the carbons, which brings them into their normal position, and the light is immediately pro-

duced. When the circuit is interrupted the armature is released; the upper ends of the carbons come into contact, and the light is produced as before. When several pairs of carbons are placed in the same circuit they are, by this arrangement, lighted simultaneously.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 24, December 9, 1878.

Report on Mr. Lawrence Smith's Memoir on the Native Iron of Greenland and the Dolerite which it Contains.—MM. Ste.-Claire Deville, Des Cloizeau, and Daubrée.—Mr. Smith concludes that all the native irons of Greenland are mutually similar, and, on the contrary, that they differ from the meteoritic irons. The former are confined to the basaltic region, extending from lat. 69° N. to 76° N., and are most abundant in its best developed regions. It is known that the terrestrial rocks which offer a resemblance to meteorites all belong to the deeper regions of the earth's crust. In those regions of Greenland where native iron occurs layers of lignites underlie here and there the great basaltic deposit, and it is possible that these carbonaceous matters may have partially reduced the iron to the metallic state. Still it is equally possible that the native iron may have been brought up from below, like the native alloy of platinum and iron which is found embedded in peridotite rocks.

Automatic Current-Regulator.—M. Hospitalier.—By means of the author's apparatus the intensity of a current may be maintained between two limits fixed beforehand, and which may be as near together as is desired.

Note on a Remarkable Specimen of Iron Silicide.—J. Lawrence Smith.—The specimen has the appearance of an ingot from the blast-furnace and has a brilliant surface. It resists nearly all reagents except hydrofluoric acid and caustic alkalis at a red heat. It contains 15 per cent of silicium, whilst the alloy of iron and manganese richest in silicium has only 10 per cent. Its origin is unknown.

A New Acid Derived from Camphor.—A. Haller.—The author has obtained a homologue of camphoric acid, which he proposes to name hydroxy-camphobic acid, $C_{11}H_{18}O_4$. It decomposes the alkaline and alkaline-earth carbonates, forming salts with their bases. The author has analysed the salts of lead, copper, and zinc.

Formation of Hexamethyl-benzin by the Decomposition of Aceton.—W. H. Greene.—The author has obtained this product in quantity by the action of aceton upon melted chloride of zinc at a high temperature.

Normal Ethyl-oxy-butyric Acid and its Derivatives.—M. Duvillier.—An examination of the ethyl-oxy-butyrate of methyl and of ethyl-oxy-butyramide.

Presence of Ytterbium in the Sipylite of Amherst (Virginia).—Marc Delafontaine.—The author has isolated from this mineral a base whose characters agree closely with those of M. Marignac's ytterbia.

Existence of Baryta and Strontia in all the Rocks forming Primordial Districts.—M. Dieulafoy.—The author finds that these earths are easily recognised in feldspars, in the mica of primitive rocks, in granites coarse or fine, and in syenite. The author considers that these rocks are the original source of baryta and strontia.

Danger of the Use of Borax for the Preservation of Food, and Causes why certain Substances Deprive

Meat of its Nutritive Properties.—G. Le Bon.—Meat steeped in a solution of pure borax, or covered with the powdered salt, may be preserved unchanged for a long time, but if taken as food such meat produces intestinal derangements, which necessitates its disuse. Borax, taken in small successive doses, is a poisonous agent, the use of which in the preservation of alimentary substances ought to be strictly prohibited. M. Peligot has already pointed out the poisonous action of borax upon plants. Several companies in America, who had begun to use this salt for the preservation of meat, have been obliged to give it up. The author further shows the necessity of avoiding saline substances altogether for the preservation of food, an object which he considers attainable solely by the use of cold.

MISCELLANEOUS.

University of London.—The following are lists of the candidates who have passed the recent examinations:—*Second B.A. and Second B.Sc. Examinations.*—*Examinations for Honours* (B.A. and B.Sc. conjointly).—*Mathematics.* First Class: J. Larmor, B.Sc. (Scholarship), St. John's College, Cambridge; W. H. Gunston,* B.A., St. John's, Cambridge. Third Class: J. A. Owen, B.Sc., private study. *Logic and Psychology.* First Class: W. K. Griffin, B.Sc., University College and private study. Second Class: S. W. Bowser, B.A., University and Regent's Park College; J. Enright, B.Sc. St. Mary's Hospital and King's College; J. Browne, B.A., Stonyhurst College. Third Class: I. Abrahams, B.A., Jews' and University Colleges; R. Gill, B.Sc., St. Bartholomew's Hospital; G. A. Stebbing, B.A., Catholic University College, Kensington; G. E. Ford, B.A., University College; J. W. Greig, B.A., University College; J. J. Beuzemaker, B.A., private study; R. A. Freeman, B.A., private study. *Classics* (B.A. only). First Class: F. C. Montague (Scholarship), Balliol College, Oxford; A. W. Lockyer, private study. Second Class: E. Etherington, Stonyhurst College; G. F. Colborne, private study and tuition; H. H. C. Thurston, Stonyhurst College; T. Slater, Stonyhurst College. Third Class: J. Browne, Stonyhurst College; C. G. Higginson, Owens College; C. T. Galton, Stonyhurst College. *French.* First Class: J. W. Greig (Prize) University College; O. E. Bodington, Giggleswick School and private tuition; F. J. Morrish, Cheshunt College and private study. Second Class: F. P. Hartley, private study; C. G. Higginson, Owens College. Third Class: E. A. Durham, private study; G. Board, private study. *German.* Second Class: C. T. Galton, Stonyhurst; J. T. Christie, King's College, London, and Exeter, Oxford. *Chemistry* (B.Sc. only). R. Gill, St. Bartholomew's Hospital; C. F. Cross, King's and Owens Colleges. *Experimental Physics.* Second Class: L. H. Edmunds, University College. Third Class: M. F. O'Reilly, St. Joseph's College, Clapham; G. W. von Tunzelmann, University College. *Physical Geography and Geology.* Second Class: T. Gough, private study and Royal College of Chemistry. Third Class: G. H. Bailey, University College and private study; P. N. Bose, University College. *Botany.* First Class: T. Gough, private study and Royal College of Chemistry. Third Class: P. N. Bose, University College. *Zoology.* Second Class: S. H. C. Martin, University College.

South London School of Pharmacy.—The sixth Annual Dinner took place at the Horns Assembly Rooms on Friday, the 20th ult. There were about 130 old and new students present, together with a large number of gentlemen interested in the progress of the School. Dr. Julius Pollock presented the prizes to the following successful students:—*Senior Chemistry.*—(Medal) Mr. Pocock; (Certificate) Miss Stammwitz. *Junior Chemistry.*—(Medal) Mr. Harrison; (Certificate) Mr. New-

* Obtained the number of marks qualifying for the Scholarship.

begin. *Botany.*—(Medal) Mr. Lord; (Certificate) Mr. Scammell. *Materia Medica.*—(Medal) Mr. Newbigin; (Certificates) Mr. Lemmon and Mr. Harrison, equal. *Practical Pharmacy and Dispensing.*—(Medal) Mr. Lemmon; (Certificate) Mr. Lord. *Silver Medal of the School, for Session 1877-78.*—Mr. John E. Phillips.

NOTES AND QUERIES.

Asphalte Pitch.—Will any of your correspondents kindly give me any information of the best method of making asphaltic pitch, and the way to make it into "blocks" ready for use?—J. R.

Peroxide of Hydrogen.—Will any of your readers inform me how I can obtain information how to make peroxide of hydrogen of ten volumes strength? I want it for commercial purposes.—THOMAS WARDLE.

Distillation of Tar.—(Reply to R. J.)—The best work on the distillation of tar, &c., is by Dr. George Lunge, published by Vieweg and Sohn, Brunswick.—DESTRUCTIVE DISTILLATOR.

MEETINGS FOR THE WEEK.

MONDAY, Jan. 6th.—Medical, 8.

London Institution, 5.

TUESDAY, 7th.—Mineralogical, at 116, Victoria Street, 8. "On Ptilolite, an unrecognised species," by Prof. M. F. Heddle. "On so-called Green Garnets from the Urals," by Prof. A. H. Church. "On the Magnetism of Rocks and Minerals," by J. B. Hannay, F.C.S. "On the Celestine and Baryto-celestine of Clifton," by J. N. Collie; communicated by W. W. Stoddart. "On some Silicates of Copper," by W. Semmons. "Contributions towards a History of British Meteorites," by Townsend M. Hall, F.G.S. "Notes on some Crystals of Iron," by Amos Beardsley, F.G.S. "Additional Note on Penwithite," by J. H. Collins, F.G.S.

Anthropological, 8.

WEDNESDAY, 8th.—Geological, 8.

Microscopical, 8.

THURSDAY, 9th.—Royal, 8.30.

FRIDAY, 10th.—Astronomical 8.

Quekett, 8.

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- III. Sanitary Science in the United States: Its Present and its Future. By Albert R. Leeds, Ph.D.
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- V. Peruvian Antiquities. By E. R. Heath, M.D.

Notices of Scientific Works, Obituary, &c.

NOTICE.

MONTHLY ISSUE OF THE JOURNAL OF SCIENCE.

The JOURNAL OF SCIENCE will in future be issued MONTHLY instead of QUARTERLY, and will consist of 48 pp., the form and general appearance remaining the same.

The first of the Monthly issue will appear on
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London: 15, Shoe Court, Ludgate Hill, E.C.

THE CHEMICAL NEWS.

VOL. XXXIX. No. 998.

DISCUSSION OF THE WORKING HYPOTHESIS THAT THE SO-CALLED ELEMENTS ARE COMPOUND BODIES.

By J. NORMAN LOCKYER, F.R.S.

(Concluded from p. 3.)

II.

*Application of the above Views to Calcium, Iron, Lithium,
and Hydrogen.*

Calcium.

was in a communication to the Royal Society made some time ago (*Proc.*, vol. xxii., p. 380, 1874), that I referred to the possibility that the well-known line-spectra of the elementary bodies might not result from vibration of similar molecules. I was led to make the

the spectra of the brighter stars before this point could be determined.

I also remarked that this result enabled us to fix with very considerable accuracy the electric dissociating conditions which are equivalent to that degree of dissociation at present at work in the sun.

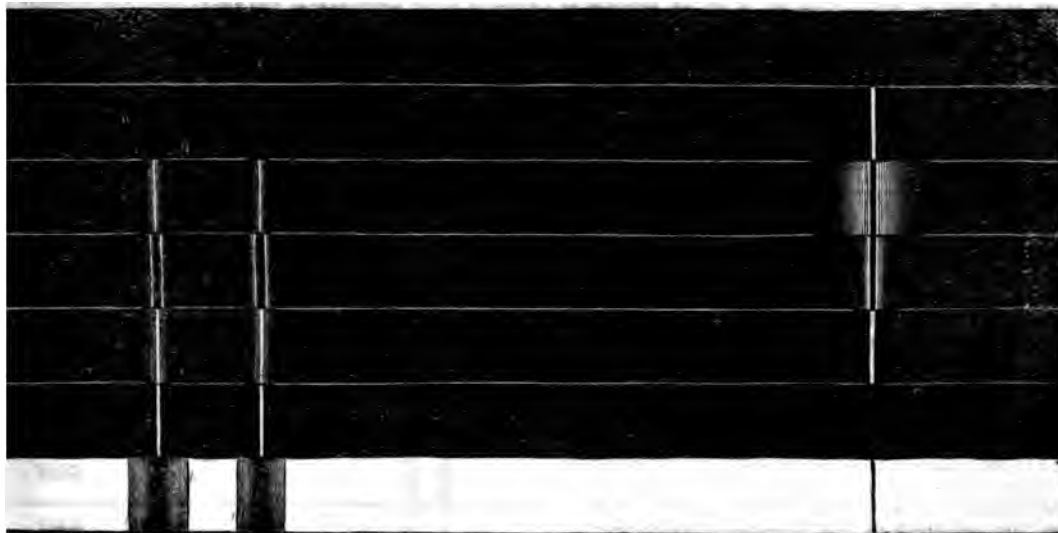
In Fig. 3 I have collected several spectra copied from photographs in order that the line of argument may be grasped.

First we see what happens to the non-dissociated and the dissociated chloride. Next we have the lines with a weak voltaic arc, the single line to the right (W. L. 4226.3) is much thicker than the two lines (W. L. 3933 and 3968) to the left, and reverses itself.

We have next calcium exposed to a current of higher tension. It will be seen that here the three lines are almost equally thick, and all reverse themselves.

Now it will be recollected, that in the case of known compounds the band structure of the true compounds is reduced as dissociation works its way, and the spectrum of each constituent element makes its appearance. If in 3 we take the wide line as representing the banded spectrum of the compound, and the thinner ones as representing the longest elemental lines making their appearance as the result of partial dissociation, we have, by hypothesis, an element behaving like a compound.

FIG. 3.—THE BLUE END OF THE SPECTRUM OF CALCIUM UNDER DIFFERENT CONDITIONS.



Calcium is combined with chlorine (CaCl_2). When the temperature is low, the compound molecule vibrates as a whole, the spectrum is at the red end, and no lines of calcium are seen. 2. The line of the metal seen when the compound molecule is dissociated to a slight extent with an induced current. 3. The spectrum of metallic calcium in the electric arc with a small number of cells. 4. The same when the number of cells is increased. 5. The spectrum when a coil and small jar are employed. 6. The spectrum when a large coil and large jar are used. 7. The absorption of the calcium vapour in the sun.

mark in consequence of the differences to which I have already drawn attention in the spectra of certain elements as observed in the spectrum of the sun and in those obtained with the ordinary instrumental appliances.

Later (*Proc. Roy. Soc.*, No. 168, 1876) I produced evidence that the molecular grouping of calcium which, with small induction-coil and small jar, gives a spectrum with its chief line in the blue, is nearly broken up in the sun, and quite broken up in the discharge from a large coil and jar, into another or others with lines in the violet.

I said "another," or "others," because I was not then able to determine whether the last-named lines proceeded from the same or different molecules; and I added that it as possible we might have to wait for photographs of

If the hypothesis be true, we ought to be able not only to obtain, with lower temperatures, a still greater preponderance of the single line, as we do; but with higher temperatures, a still greater preponderance of the double ones, as we do.

I tested this in the following manner:—Employing photography, because the visibility of the more refrangible lines is small, and because a permanent record of an experiment, free as it must be from all bias, is a very precious thing.

Induced currents of electricity were employed in order that all the photographic results might be comparable.

To represent the lowest temperature I used a small induction coil and a Leyden jar only just large enough to secure the requisite amount of photographic effect. To represent the highest, I used the largest coil and jar at my disposal. The spark was then taken between two alu-

* Paper read at the Royal Society, December 12, 1878.

minium electrodes, the lower one cup-shaped, and charged with a salt of calcium.

In the figure I give exact copies of the results obtained. It will be seen that with the lowest temperature only the single line (2) and with the highest temperature only the two more refrangible lines (6) are recorded on the plate.

This proved that the intensity of the vibrations was quite changed in the two experiments.

Perhaps it may not be superfluous here to state the reasons which induced me to search for further evidence in the stars.

It is abundantly clear that if the so-called elements, or, more properly speaking, their finest atoms—those that give us line spectra—are really compounds, the compounds must have been formed at a very high temperature. It is easy to imagine that there may be no superior limit to temperature, and therefore no superior limit beyond which such combinations are possible, because the atoms which have the power of combining together at these transcendental stages of heat do not exist as such, or rather they exist combined with other atoms, like or unlike, at all lower temperatures. Hence association will be a combination of more complex molecules as temperature is reduced, and of dissociation, therefore, with increased temperature there may be no end.

That is the first point.

The second is this:—

We are justified in supposing that our "calcium," once formed, is a distinct entity whether it be an element or not, and therefore, by working at it alone, we should never know whether the temperature produces a single simpler form or more atomic condition of the same thing, or whether we actually break it up into $X + Y$, because neither X nor Y will ever vary.

But if calcium be a product of a condition of relatively lower temperature, then in the stars, hot enough to enable its constituents to exist uncombined, we may expect these constituents to vary in quantity; there may be more of X in one star and more of Y in another; and if this be so, then the H and K lines will vary in thickness, and the extremest limit of variation will be that we shall only have H representing, say X in one star, and only have K representing say Y in another. Intermediately between these extreme conditions we may have cases in which, though both H and K are visible, H is thicker in some and K is thicker in others.

Prof. Stokes was good enough to add largely to the value of my paper as it appeared in the *Proceedings* by appending a note pointing out that "When a solid body such as a platinum wire, traversed by a voltaic current, is heated to incandescence, we know that as the temperature increases not only does the radiation of each particular refrangibility absolutely increase, but the proportion of the radiations of the different refrangibilities is changed, the proportion of the higher to the lower increasing with the temperature. It would be in accordance with analogy to suppose that as a rule the same would take place in an incandescent surface, though in this case the spectrum would be discontinuous instead of continuous. Thus if A, B, C, D, E denote conspicuous bright lines of increasing refrangibility, in the spectrum of the vapour, it might very well be that at a comparatively low temperature A should be the brightest and the most persistent; at a higher temperature, while all were brighter than before, the relative brightness might be changed, and C might be the brightest and the most persistent, and at a still higher temperature E ."

On these grounds Prof. Stokes, while he regarded the facts I mentioned as evidence of the high temperature of the sun, did not look upon them as *conclusive* evidence of the dissociation of the molecule of calcium.

Since that paper was sent in, however, the appeal to the stars to which I referred in it has been made, and made with the most admirable results, by Dr. Huggins.

The result of that appeal is that the line which, according to Prof. Stokes's view, should have prevailed over all

others, as Sirius is acknowledged to be a hotter star than our sun, is that if it exists at all in the spectrum, it is so faint that it was recognised by Dr. Huggins in the first instance.

In Sirius, indeed, the H line due to one molecular grouping of calcium is as thick as are the hydrogen lines as mapped by Secchi, while the K line, due to another molecular grouping, which is equally thick in the spectrum of the sun, has not yet made its appearance.

In the sun, where it is as thick as H , the hydrogen lines have vastly thinned.

While this paper has been in preparation, Dr. Huggins has been good enough to communicate to me the results of his most important observations, and I have also had an opportunity of inspecting several of the photographs which he has recently taken. The result of the recent work has been to show that H and K are of about the same breadth in Sirius. In α Aquilæ while the relation of H to K is not greatly changed, a distinct approach to the solar condition is observed, K being now unmistakably present, although its breadth is small as compared with that of H . I must express my obligations to Dr. Huggins for granting me permission to enrich my paper with reference to these unpublished observations. His letter, which I have permission to quote, is as follows:—

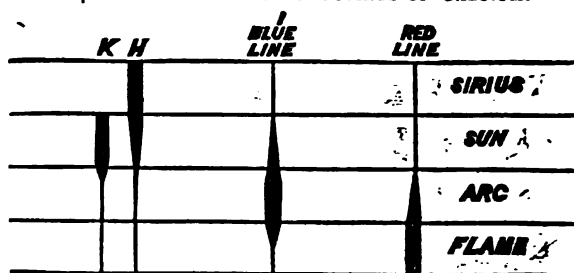
"It may be gratifying to you to learn that in a photograph I have recently taken of the spectrum of α Aquilæ there is a line corresponding to the more refrangible of the solar H lines [that is K], but about half the breadth of the line corresponding to the first H lines.

"In the spectra of α Lyrae and Sirius the second line is absent."

Prof. Young's observations of the chromospheric lines, to which I shall afterwards refer, give important evidence regarding the presence of calcium in the chromosphere of the sun. He finds that the H and K lines of calcium are strongly reversed in every important spot, and that in solar storms H has been observed injected into the chromosphere seventy-five times, and K fifty times, while the blue line at $W. L. 4226.3$, the all-important line at the arc-temperature, was only injected thrice.

Further, in the eclipse observed in Siam in 1875, the H and K lines left the strongest record in the spectrum of

FIG. 4.—THE MOLECULAR GROUPINGS OF CALCIUM.



the chromosphere, while the line near G in a photographic region of much greater intensity was not recorded at all. In the American eclipse of the present year the H and K lines of calcium were distinctly visible at the base of the corona, in which for the first time the observers could scarcely trace the existence of any hydrogen.

To sum up, then, the facts regarding calcium, we have first of all the H -line differentiated from the others by its almost solitary existence in Sirius. We have the K -line differentiated from the rest by its birth, so to speak, in α Aquilæ, and the thickness of its line in the sun, as compared to that in the arc. We have the blue line differentiated from H and K by its thinness in the solar spectrum while they are thick, and by its thickness in the arc while they are thin. We have it again differentiated from them by its absence in solar storms in which they are almost universally seen, and finally, by its absence during eclipses,

while the H and K lines have been the brightest seen or photographed. Last stage of all, we have calcium, distinguished from its salts by the fact that the blue line is only visible when a high temperature is employed, each salt having a definite spectrum of its own, in which none of the lines to which I have drawn attention appear, so long as the temperature is kept below a certain point.

Iron.

With regard to the iron spectrum I shall limit my remarks to that portion of it visible on my photographic plates between H and G. It may be described as a very complicated spectrum, so far as the number of lines is concerned in comparison with such bodies as sodium and potassium, lead, thallium, and the like, but unlike them again it contains no one line which is clearly and unmistakably reversed on all occasions. Compared, however, with the spectrum of such bodies as cerium and uranium the spectrum is simplicity itself.

Now among these lines are two triplets, two sets of three lines each, giving us beautiful examples of those repetitions of structure in the spectrum which we meet with in the spectra of almost all bodies, some of which have already been pointed out by Mascart, Cornu, and myself. Now the facts indicate that these two triplets are not due to the vibration of the same molecular grouping which gives rise to most of the other lines. They are as follows:—In many photographs in which iron has been compared with other bodies, and in others again in which iron has been photographed as existing in different degrees of impurity in other bodies, these triplets have been seen almost alone, and the relative intensity of them, as compared with the few remaining lines, is greatly changed. In this these photographs resemble one I took three years ago, in which a large coil and jar were employed instead of the arc, which necessitated an exposure of an hour instead of two minutes. In this the triplet near G is very marked, the two adjacent lines more refrangible near it, which are seen nearly as strong as the triplet itself in some of the arc photographs I possess, are only very faintly visible, while dimmer still are seen the lines of the triplet between H and k.

There is another series of facts in another line of work. In solar storms, as is well known, the iron lines sometimes make their appearance in the chromosphere. Now if we were dealing here with one molecular grouping, we should expect the lines to make their appearance in the order of their lengths, and we should expect the shortest lines to occur less frequently than the longest ones. Now, precisely the opposite is the fact. One of the most valuable contributions to solar physics that we possess is the memoir in which Prof. C. A. Young records his observation of the chromospheric lines, made on behalf of the United States Government, at Sherman, in the Rocky Mountains. The glorious climate and pure air of this region, to which I can personally testify, enabled him to record phenomena which it is hopeless to expect to see under less favourable conditions. Among these were injections of iron vapour into the chromosphere, the record taking the form of the number of times any one line was seen during the whole period of observation.

Now two very faint and short lines close to the triplet near G were observed to be injected thirty times, while one of the lines of the triplet was only injected twice.

The question next arises, are the triplets produced by one molecular grouping or by two? This question I also think the facts help us to answer. I will first state by way of reminder that in the spark photograph the more refrangible triplet is barely visible, while the one near G is very strong. Now if one molecular grouping alone were in question this relative intensity would always be preserved however much the absolute intensity of the compound system might vary, but if it is a question of two molecules we might expect that in some of the regions open to our observation we should get evidence of cases in which the relative intensity is reversed or the

two intensities are assimilated. What might happen does happen; the relative intensity of the two triplets in the spark photograph is grandly reversed in the spectrum of the sun. The lines barely visible in the spark photograph are among the most prominent in the solar spectrum, while the triplet which is strong in that photograph is represented by Fraunhofer lines not half so thick. Indeed, while the hypothesis that the iron lines in the region I have indicated are produced by the vibration of one molecule does not include all the facts, the hypothesis that the vibrations are produced by at least three distinct molecules includes all the phenomena in a most satisfactory manner.

Lithium.

Before the maps of the long and short lines of some of the chemical elements compared with the solar spectra, which were published in the *Phil. Trans.* for 1873, "Plate IX.," were communicated to the Society, I very carefully tested the work of prior observers on the non-coincidence of the red and orange lines of that metal with the Fraunhofer lines, and found that neither of them were strongly if at all represented in the sun, and this remark also applies to a line in the blue at wave-length 4603.

The photographic lithium line, however, in the violet has a strong representative among the Fraunhofer lines.

Applying, therefore, the previous method of stating the facts, the presence of this line in the sun differentiates it from all the others. For the differentiation of the red and yellow lines I need only refer to Bunsen's spectral analytical researches, which were translated in the *Phil. Mag.*, December, 1875.

In Plate IV. two spectra of the chloride of lithium are given, one of them showing the red line strong and the yellow one feeble, the other showing merely a trace of the red line, while the intensity of the yellow one is much increased, and a line in the blue is indicated. Another notice of the blue line of lithium occurs in a discourse by Prof. Tyndall, reprinted in the *CHEMICAL NEWS*, and a letter of Dr. Frankland's to Prof. Tyndall, dated November 7, 1861. This letter is so important for my argument that I reprint it entire from the *Phil. Mag.*, vol. xxii., p. 472:—

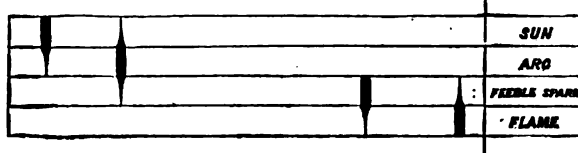
"On throwing the spectrum of lithium on the screen yesterday, I was surprised to see a magnificent blue band. At first I thought the lithic chloride must be adulterated with strontium, but on testing it with Steinheil's apparatus it yielded normal results without any trace of a blue band. I am just now reading the report of your discourse in the *CHEMICAL NEWS*, and I find that you have noticed the same thing. Whence does this blue line arise? Does it really belong to the lithium, or are the carbon points or ignited air guilty of its production? I find there blue bands with common salt, but they have neither the definiteness nor the brilliancy of the lithium band. When lithium wire burns in air it emits a somewhat crimson light; plunge it into oxygen, and the light changes to bluish white. This seems to indicate that a high temperature is necessary to bring out the blue ray."

"POSTSCRIPT, Nov. 22, 1861.—I have just made some further experiments on the lithium spectrum, and they conclusively prove that the appearance of the blue line depends entirely on the temperature. The spectrum of lithic chloride, ignited in a Bunsen's burner flame, does not disclose the faintest trace of the blue line; replace the Bunsen's burner by a jet of hydrogen (the temperature of which is higher than that of the Bunsen's burner) and the blue line appears faint, it is true, but sharp and quite unmistakable. If oxygen now be slowly turned into the jet, the brilliancy of the blue line increases until the temperature of the flame rises high enough to fuse the platinum, and thus put an end to the experiment."

These observations of Profs. Tyndall and Frankland differentiate this blue line from those which are observed

at low temperatures. The line in the violet to which I have already referred is again differentiated from all the rest by the fact that it is the only line in the spectrum of the sun which is strongly reversed, so far as our present knowledge extends. The various forms of lithium, therefore, may be shown in the following manner.

FIG. 5.—THE MOLECULAR GROUPINGS OF LITHIUM.



It is remarkable that in the case of this body which at relatively low temperature goes through its changes, its compounds are broken up at the temperature of the Bunsen burner. The spectrum, *e.g.* of the chloride, so far as I know, has never been seen.

Hydrogen.

All the phenomena of variability and inversion in the order of intensity presented to us in the case of calcium can be paralleled by reference to the knowledge already acquired regarding the spectrum of hydrogen.

Dr. Frankland and myself were working together on the subject in 1869. In that year (*Proc.*, No. 112) we pointed out that the behaviour of the *k* line was *hors ligne*, and that the whole spectrum could be reduced to one line, F.

"1. The Fraunhofer line on the solar spectrum, named *k* by Ångström, which is due to the absorption of hydrogen, is not visible in the tubes we employ with low battery and Leyden-jar power; it may be looked upon, therefore, as an indication of relatively high temperature. As the line in question has been reversed by one of us in the spectrum of the chromosphere, it follows that the chromosphere, when cool enough to absorb, is still of a relatively high temperature.

"2. Under certain conditions of temperature and pressure, the very complicated spectrum of hydrogen is reduced in our instrument to one line in the green corresponding to F in the solar spectrum."

As in the case of calcium also, solar observation affords us most precious knowledge. The *k* line was missing from the protuberances in 1875, as will be shown from the accompanying extract from the Report of the Eclipse Expedition of that year:—

"During the first part of the eclipse two strong protuberances close together are noticed; on the limb towards the end these are partially covered, while a series of protuberances came out at the other edge. The strongest of these protuberances are repeated three times, an effect of course of the prism, and we shall have to decide if possible the wave-lengths corresponding to the images. We expect *a priori* to find the hydrogen lines represented. We know three photographic hydrogen lines: F, a line near G, and *k*. F is just at the limit of the photographic part of the spectrum, and we find indeed images of protuberances towards the less refrangible part at the limit of photographic effect. For, as we shall show, a continuous spectrum in the lower parts of the corona has been recorded, and the extent of this continuous spectrum gives us an idea of the part of the spectrum in which each protuberance line is placed. We are justified in assuming, therefore, as a preliminary hypothesis, that the least refrangible line in the protuberance shown on the photograph is due to F, and we shall find support of this view in the other lines. In order to determine the position of the next line the dispersive power of the prism was investigated. The prism was placed on a goniometer table in minimum deviation for F, and the angular distance between F and the hydrogen line near G, *i.e.*, Hy,

was found, as a mean of several measurements to be 3'. The goniometer was graduated to 15", and owing to the small dispersive power, and therefore relatively great breadth of the slit, the measurement can only be regarded as a first approximation. Turning now again to our photographs, and calculating the angular distance between the first and second ring of protuberances, we find that distance to be 3' 15". We conclude, therefore, that this second ring is due to hydrogen. We, therefore, naturally looked for the third photographic hydrogen line, which is generally called *k*, but we found no protuberance on our photographs corresponding to that wave-length. Although this line is always weaker than Hy, its absence on the photograph is rather surprising, if it be due to the fact that the line is one which only comes out at a high temperature. This is rendered likely by the researches of Frankland and Lockyer (*Proc. Roy. Soc.*, vol. xvii., p. 453).

"We now turn to the last and strongest series of protuberances shown on our photographs. The distance between this series and the one we have found reason for identifying with Hy is very little greater than that between H β and Hy. Assuming the distances equal, we conclude that the squares of the inverse wave-lengths of the three series are in arithmetical progression. This is true as a first approximation. We then calculated the wave-length of this unknown line, and found it to be approximately somewhat smaller than 3957 tenth-metres. No great reliance can be placed, of course, on the number, but it appears that the line must be close to the end of the visible spectrum.

"In order to decide if possible what this line is due to, we endeavoured to find out both by photography and fluorescence whether hydrogen possesses a line in that part of the spectrum. We have not at present come to any definite conclusion. In vacuum tubes prepared by Geissler containing hydrogen, a strong line more refrangible than H is seen, but these same tubes show between Hy and H δ , other lines known not to belong to hydrogen, and the origin of the ultra violet line is therefore difficult to make out. We have taken the spark in hydrogen at atmospheric pressures, as impurities are easier to eliminate, but a continuous spectrum extends over the violet and part of the ultra-violet, and prevents any observation as to lines. We are going on with experiments to settle this point.

"Should it turn out that the line is not due to hydrogen, the question will arise what substance it is due to. It is a remarkable fact that the calculated wave-length comes very close to H. Young has found that these calcium lines are always reversed in the penumbra and immediate neighbourhood of every important sun-spot, and calcium must therefore go up high into the chromosphere. We draw attention to this coincidence, but our photographs do not allow us to draw any certain conclusions.

"At any rate, it seems made out by our photographs that the photographic light of the protuberances is in great part due to an ultra-violet line which does not certainly belong to hydrogen. The protuberances as photographed by this ultra-violet ray seem to go up higher than the hydrogen protuberances, but this may be due to the relative greater length of the line."

In my remarks upon calcium I have already referred to the fact that the line which our observation led us to believe was due to calcium in 1875, was traced to that element in this year's eclipse. The observations also show the curious connection that, at the time when the hydrogen lines were most brilliant in the corona, the calcium lines were not detected; next, when the hydrogen lines, being still brilliant, the *k* line was not present (a condition of things which, in all probability, indicated a reduction of temperature), calcium began to make itself unmistakably visible; and finally, when the hydrogen lines are absent, H and K become striking objects in the spectrum of the corona.

To come back to *k* then, I have shown that Dr. Frank-

land and myself, in 1869, found that it only made its appearance when a high tension was employed. We have seen that it was absent from among the hydrogen lines during the eclipse of 1875.

I have now to strengthen this evidence by the remark that it is always the shortest line of hydrogen in the chromosphere.

I now pass to another line of evidence.

I submit to the Society a photograph of the spectrum of indium, in which, as already recorded by Thalén, the strongest line is one of the lines of hydrogen (*h*), the other line of hydrogen (near *G*) being absent. I have observed the *C* line in the spark produced by the passage of an induced current between indium poles in dry air.

As I am aware how almost impossible it is to render air perfectly dry, I made the following differential experiment. A glass tube with two platinum poles about half an inch apart was employed. Through this tube a slow current of air was driven after passing through a U-tube one foot high, containing calcic chloride, and then through sulphuric acid in a Wolff's bottle. The spectrum of the spark passing between the platinum electrodes was then observed, a coil with five Grove cells and a medium-sized jar being employed. Careful notes were made of the brilliancy and thickness of the hydrogen lines as compared with those of air. This done, a piece of metallic indium which was placed loose in the tube, was shaken so that one part of it rested against the base of one of the poles, and one of its ends at a distance of a little less than half an inch from the base of the other pole. The spark then passed between the indium and the platinum. The red and blue lines of hydrogen were then observed both by my friend, Mr. G. W. Hemming, Q.C., and myself. Their brilliancy was most markedly increased. This unmistakable indication of the presence of hydrogen, or rather of that form of hydrogen which gives us the *h* line alone associated into that form which gives us the blue and red lines, showed us that in the photograph we were not dealing with a physical coincidence, but that in the arc this special form of hydrogen had really been present; that it had come from the indium, and that it had registered itself on the photographic plate, although ordinary hydrogen persistently refuses to do so. Although I was satisfied from former experiments that occluded hydrogen behaves in this respect like ordinary hydrogen. I begged my friend, Mr. W. C. Roberts, F.R.S., chemist to the Mint, to charge a piece of palladium with hydrogen for me. This he at once did, and I take this present opportunity to express my obligation to him. I exhibit to the Society a photograph of this palladium and of indium side by side. It will be seen that one form of hydrogen in indium has distinctly recorded itself on the plate, while that in palladium has not left a trace. I should add that the palladium was kept in a sealed tube till the moment of making the experiment, and that special precautions were taken to prevent the two pieces between which the arc was taken becoming unduly heated.

To sum up, then, the facts with regard to hydrogen; we have *h* differentiated from the other lines by its appearance alone in indium; by its absence during the eclipse of 1875, when the other lines were photographed; by its existence as a short line only in the chromosphere of the sun, and by the fact that in the experiments of 1869 a very high temperature was needed to cause it to make its appearance.

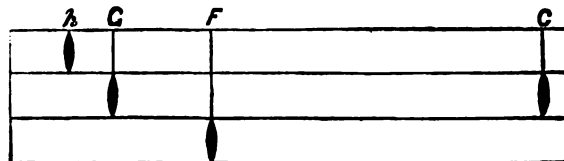
With regard to the isolation of the *F* line I have already referred to other experiments in 1869, in which Dr. Frankland and myself got it alone.* I exhibit to the Society a globe containing hydrogen which gives us the *F* line without either the red or the blue one.

The accompanying drawing shows how these lines are integrated in the spectrum of the sun.

I have other evidence which, if confirmed, leads to the conclusion that the substance which gives the non-

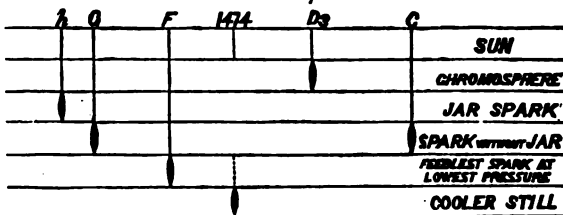
reversed line in the chromosphere and the line at 1474 of Kirchhoff's scale, termed the coronal line, are really other forms of hydrogen. One of these is possibly more simple than that which gives us *h* alone, the other more complex than that which gives us *F* alone. The evidence on this point is of such extreme importance to solar physics, and throws so much light on star structure generally, that I am now engaged in discussing it and shall reserve it for a special communication.

FIG. 6.



In the meantime I content myself by giving a diagram in which I have arranged the various groupings of hydrogen as they appear to exist, from the regions of highest to those of lowest temperature in our central luminary.

FIG. 7.



Summation of the above Series of Facts.

I submit that the facts above recorded are easily grouped together, and a perfect continuity of phenomena established on the hypothesis of successive dissociations analogous to those observed in the cases of undoubted compounds.

The other Branches of the Inquiry.

When we pass to the other possible evolutionary processes to which I have before referred, and which I hope to discuss on a future occasion, the inquiry becomes much more complicated by the extreme difficulty of obtaining pure specimens to work with, although I should remark that in the working hypothesis now under discussion the cause of the constant occurrence of the same substance as an impurity in the same connection is not far to seek. I take this opportunity of expressing my obligations to many friends who have put themselves to great trouble in obtaining specimens of pure chemicals for me during the whole continuance of my researches. Among these I must mention Dr. Russell, who has given me many specimens prepared by the lamented Matthiessen, as well as some of cobalt and nickel prepared by himself; Prof. Roscoe, who has supplied me with vanadium and caesium alum; Mr. Crookes, who has always responded to my call for thallium; Mr. Roberts, chemist to the Mint, who has supplied me with portions of the gold and silver trial plates and some pieces of palladium; Dr. Hugo Müller, who has furnished me with a large supply of electrolytically-deposited copper; Mr. Holtzman, who has provided me with cerium, lanthanum, and didymium prepared by himself; Mr. George Matthey, of the well-known metallurgical firm of Johnson and Matthey, who has provided me with magnesium and aluminium of marvellous purity; while to Mr. Valentin, Mr. Mellor, of Salford, and other friends, my thanks are due for other substances.

I have already pointed out that a large portion of the work done in the last four years has consisted in the elimination of the effects of impurities. I am therefore aware

* See also Plücker, *Phil. Trans.*, 1865, part 1, p. 21.

of the great necessity for caution in the spectroscopic examination of various substances. There is, however, a number of bodies which permit of the inquiry into their simple or complex nature being made in such a manner that the presence of impurities will be to a certain extent negligible. I have brought this subject before the Royal Society at its present stage, in the hope that possibly others may be induced to aid enquiry in a region in which the work of one individual is as a drop in the ocean. If there is anything in what I have said, the spectra of all the elementary substances will require to be re-mapped and re-mapped from a new standpoint; further, the arc must replace the spark, the photography must replace the eye. A glance at the red end of the spectrum of almost any substance incandescent in the voltaic arc in a spectroscopic of large dispersion, and a glance at the maps prepared by such eminent observers as Huggins and Thalén, who have used the coil, will give an idea of the mass of facts which have yet to be recorded and reduced before much further progress can be made.

In conclusion, I would state that only a small part of the work to which I have drawn attention is my own. In some cases I have merely, as it were, codified the work done by other observers in other countries. With reference to that done in my own laboratory, I may here repeat what I have said before on other occasions, that it is largely due to the skill, patience, and untiring zeal of those who have assisted me. The burthen of the final reduction, to which I have before referred, has fallen to Mr. Miller, my present assistant; while the mapping of the positions and intensities of the lines was done by Messrs. Friswell, Meldola, Ord, and Starling, who have successfully filled that post.

I have to thank Corporal Ewings, R.E., for preparing the various diagrams which I have submitted to the notice of this Society.

ON SUPERSATURATED SOLUTIONS.

By J. G. GRENFELL.

In September, 1877, Professor Tomlinson read a paper before the Royal Society, the main object of which was to upset the conclusions to which I had been led by my experiments on supersaturated solutions, and thus to maintain his theory that the salt adheres to a film of any greasy substance, while the water does not, and thus separation ensues.

Before answering this paper I wished to repeat some of my experiments, and this I have not been able to do till quite recently. Those conclusions of mine which are openly or implicitly attached in the paper are as follows:—

1. Supersaturated solutions can in many cases be exposed to the air for an indefinite time, can be rubbed with dust or oil and yet not crystallise. At a meeting of the Chemical and Physical Section of the Bristol Naturalists' Society I treated strong solutions of sodium sulphate and acetate in this way, and a good many persons present did the same. I used ordinary castor oil.
2. Absorbent substances, when not at once saturated by the solution, and provided the absorption is not too rapid, cause crystallisation, sometimes as the normal salt, sometimes as a modified salt. For instance, I recently put a considerable quantity of a 6 to 1 solution of sodium carbonate on a new sheet of thick blotting-paper. In about a minute it gave a cake of the modified 7-atom salt. The crystals were quite inactive in a 2 to 1 solution. At the same time I put 13 small drops round the edge; of these two crystallised at once as the 7-atom salt, and they and the central mass were soon quite dry; the rest remained liquid exposed to air

for many hours. In this case the rapidity of the absorption prevented the formation of crystals. Swedish filter-paper generally gives the normal salt in the centre with this solution, whilst any number of small drops are absorbed, and a great many little bits of the paper can be put into drops or into the flask, and are inactive because saturated at once.* The absorbents seem to act by abstracting water. This explains the action of aerial nuclei; they act by causing gentle absorption, which gives the normal salt, or possibly by gentle evaporation. There is some reason to believe that gentle evaporation will give the normal salt just as the more rapid evaporation of drops exposed to air gives a salt with less water. Salts vary very much in their sensitiveness to absorbents, sodium sulphate being the most so that I have yet tried.

3. I believe that most cases of sudden crystallisation on removing cotton-wool, or piercing a bladder, or on giving a jerk, as Professor Tomlinson does to flatten a lens of oil into a film, or (as has often happened to me) on simply entering the room, may be explained thus:—After heating a solution convection currents of air may be seen rising and carrying drops of the solution. These settle on the cover or round the mouth, crystallise there by absorption or evaporation, and are shaken down by any sudden jar. When the covers are loose, or are gently removed, crystallisation seldom follows.

Turning now to Prof. Tomlinson's paper, I find that he has repeated a certain number of my experiments, with the result, as he believes, of upsetting my theories. As a matter of fact, I have to thank him for a very pretty set of illustrations of them.

With regard to the first point, for instance—that of exposure—he finds as I did that drops of sodium acetate can be exposed for weeks, the carbonate for days, the sulphate for many hours on paper and other surfaces. He makes a great point of the fact that this happens only in damp weather; in dry weather he says the drops crystallise. Nearly half of his paper is devoted to experiments which prove this statement. He does not, however, attempt to explain on his own theory why this should be the case, but remarks that laborious investigation is needed to determine the conditions under which bodies do or do not act as nuclei. I could not have wished for a neater proof that my theory is correct. If the nuclei act by absorption and evaporation it is obvious that they will have much less effect in wet weather than in dry. I could have predicted the result, though I have never paid special attention to the point.

I should like to know, however, what Prof. Tomlinson means by saying that drops crystallise in dry weather. There is some reason to believe that he was satisfied with seeing crystals after the lapse of a certain time, and did not try to find out what the crystals were.

This certainly seems to be the case in one instance. He says that potash alum crystallised into an opaque white mass or into transparent crystals in concentric circles round a central pit. Now the alums generally evaporate on glass, giving a more or less transparent crust with a central depression, underneath which little botryoidal masses are often formed. It is not at all common for alum to crystallise in concentric rings of separate crystals, and if, as I suppose, Prof. Tomlinson refers to the ordinary crust, he would find that this is not alum at all, as it is quite inactive in drops, while the merest trace of the normal converts it into the opaque white normal alum.

Be this, however, as it may, it is not hard to account for the statement that in dry weather the drops crystallise rapidly. It is impossible to succeed in dry weather with-

* These experiments so were shown at the meeting mentioned above.

out taking certain precautions. The rough glass plate must have been recently washed, so as to saturate with moisture the absorbent substances which exist in the depressions of the glass, and which are not rubbed off when the surface is wiped dry. If these are allowed to dry thoroughly they start crystallisation in the depressions of the plate; the crystals grow sometimes very rapidly, sometimes very slowly. In the latter case scratching with a sharp point brings the crystals into contact with the solution, which solidifies at once.

This may account for a good many of Prof. Tomlinson's negative results. It also explains his statement that drawing a sharp edge across the drops generally makes them crystallise. If the plate is in the right condition, they do not do so. I have drawn sharp points across hundreds of drops of the strongest solutions in all kinds of weather without causing crystallisation.

It is also necessary in the case of the alums or sodium acetate to clean the sharp point after every experiment with boiling water or preferably a red heat. Cold water and wiping will not generally clean a penknife for instance, owing to the great adhesiveness of the salt.

Next, with regard to rubbing with oil, which is really the crucial test for Prof. Tomlinson's theory, he seems to have been satisfied with rubbing the sulphate with oil on a glass plate; he very naturally found that it crystallised. I had extreme difficulty at first in getting this experiment to succeed, and I could not do so at all in my laboratory, where the dust possibly contains sodium sulphate. But I did succeed completely in another room.

However, if Prof. Tomlinson will wash his hands he may do what I have just done, *i.e.*, rub a little castor oil on the palm of the hand, and then rub in drops of a 3 to 1 solution of the sulphate without its crystallising. He may repeat this with sodium acetate of any strength. Strong sodium carbonate will give a modified salt which is quite inactive in a 6 to 1 solution, and is, therefore, neither the normal nor 7-atom salt. I have succeeded with the alums also, but they are difficult to manage.

Secondly, as direct evidence against the absorption theory, Prof. Tomlinson gives the following experiments: Three pieces of dried sponge were introduced into flasks of the sulphate, and were inactive. Of course they were, being saturated at once. Curiously enough, he actually describes the converse of this experiment, which is needed to maintain my theory. He found some years ago that when sponge is drawn rapidly over the surface, the solution crystallises on the sponge, while that in the flask did not. The only inference he draws from this is that results are often contradictory.

He mentions one more fact as bearing on this point. A solution of the sulphate crystallised when poured on the soil of his garden. This is the very same experiment which first led me to work at the absorption theory. I found, as described in *Nature*, that lumps of earth could be put freely into the flask; but I also found that every single drop put on the earth in the beds crystallised at once.

Similarly with regard to the third point, the effect of sudden jars, Prof. Tomlinson gives experiments which simply confirm my results. He begins thus: "There is one point upon which all observers down to the time of Mr. Grenfell are agreed, namely, that if a supersaturated solution boiling in a flask be tied over with bladder and left until cold, it will, if the bladder be pierced, immediately become solid. Indeed, this has long been a common lecture-room experiment. But Mr. Grenfell goes far beyond anything I or previous observers have done; he exposes these solutions, &c." He thus distinctly implies that I have denied the fact. Of course I have never denied it at all. In my first paper in the *Proc. Roy. Soc.*, two years ago, I carefully explained that I had been driven to abandon cotton-wool, which was generally recommended as a covering, because I felt convinced that in removing it I shook down some crystals. I added that by using loose paper covers I found that the solutions very

seldom crystallised when uncovered. Having thus saddled me with an opinion I have almost expressly repudiated, Prof. Tomlinson proceeds to upset me by the following evidence. He took two flasks of the sulphate; pierced the bladder of one, and it crystallised; he removed the cover of the other in the country, and it remained liquid during a long walk. This, as usual, supports my views. Removing the cover quietly is a very different process to piercing with a pin, and no crystals were shaken down. In the same way he shook a solution of alum several times without shaking down any crystals, but piercing with a pin was effectual. Alum is an adhesive salt, and requires a jerk to be given. Again he pierced the bladder of two flasks of the sulphate, and only one crystallised. On increasing the opening next day to the size of a pea, the other solution crystallised in a few hours. Possibly an absorbent nucleus got in through the enlarged opening, but it is more probable that the crystals round the opening effloresced and some fragment fell in. I have found the effloresced sulphate invariably active. In air it never gets quite free from the normal salt.

To sum up: the great majority of the experiments described in this paper confirm my theories. Prof. Tomlinson arrived at negative results in other cases from not knowing the precautions which have to be taken.

The fact that paper, earth, and all kinds of substances can be introduced into the flasks; that drops will remain liquid on all kinds of surfaces for hours; even granting, which I do not, that it is only in wet weather, and the experiments with oil are absolutely fatal to any theory based on preferential adhesion of the salt to a greasy surface.

The immediate production of the modified seven atom carbonate by blotting-paper, the experiments with sponge and earth, the effect of damp weather on aerial nuclei, and a multitude of similar results which I could give if necessary, are incapable of explanation on any other theory than that of the abstraction of water by absorbent bodies.

The old theory which made the air a store-house of all kinds of crystals is dead and buried. I frequently have half a dozen plates covered with drops of various solutions exposed to air in my laboratory for hours or days, and the drops evaporate quietly as modified salts, or remain liquid. The minutest trace of a crystal makes them crystallise at once.

UPON THE ALTERATION OF STANDARD AMMONIUM CHLORIDE SOLUTION WHEN KEPT IN THE DARK.

By ALBERT R. LEEDS, Ph.D.

In February, 1877, MM. Schloesing and Muntz communicated to the French Academy (*Comptes Rendus*, lxxiv., 301), the results of elaborate experiments tending to prove that nitrification was due in certain cases to the action of an organised ferment. They further showed that nitrification could be arrested in such cases by the vapour of chloroform, and would not set up again until after the filtered sewage employed had been freshly "seeded" by the washings of a soil which was known to nitrify. These observations were verified and extended by R. Warington (*CHEM. NEWS*, xxxvi., 262, 1877), who showed that chloroform and carbon disulphide effectually prevent nitrification in certain cases, while carbolic acid is probably effective to the extent in which it comes in contact with the soil. Moreover that the action of the nitrifying body does not occur at all in the light, darkness being apparently essential. In a note upon the ferment theory of nitrification, by Prof. F. H. Storer (*CHEM. NEWS*, xxxvii., 268), he states that a solution of ammonium chloride, which had been prepared ten or twelve months

previously, gave a strong reaction for nitrites. It had been preserved in a half-filled glass-stoppered bottle, put away in a dark cupboard. Similar solutions, seeded in various ways, but left in a strong light for a number of months, gave no reaction whatever for nitrites or nitrates. The following experiments confirm these results, so far as the change of the ammonium solutions in the dark is concerned, and give the rate at which it may be expected to occur under ordinary circumstances.

A standard solution of ammonium chloride had been prepared October 15th, 1877, by dissolving 0.7867 grm. NH_4Cl , purified by repeated sublimation, in a litre of redistilled water, free from ammonia. Another solution was prepared by diluting 100 c.c. of this water to 1 litre, and used constantly during the months of October and November, 1877, in making the estimations which will be found recorded in an article entitled "Contributions from the Laboratory of the Stevens Institute of Technology." They were put aside in a dark closet, and remained there until this autumn, when a new standard ammonium chloride solution was prepared from the fear that the old solutions had altered. This proves to have been the case. A careful determination, as ammonio-platinic chloride of the ammonia contained in 250 c.c. of the stronger solution, gave 0.0575 grm. or 0.00023 grm. in 1 c.c., instead of 0.00025, the amount which the solution originally contained. Instead of being perfectly limpid, as it was when made, the solution contained a number of white filaments, looking like the vestiges of some vegetable growth. The more dilute solution presented the same appearances, but not to so marked a degree. 25 c.c. of the solution were used in the examination for nitrites, but their presence could not be shown in this large quantity of the liquid. 5 c.c. of the solution were evaporated three times to dryness in a platinum dish with 0.1 grm. of caustic soda, prepared from sodium, and which had been found entirely free from nitrogen compounds. It was then distilled with 6 grms. pig-iron in a vessel entirely made of glass, the pig-iron having been previously repeatedly distilled with water, until it and the vessel had ceased to give any reaction for ammonia. The distillate contained 0.03 m.grm. H_2N equivalent to 0.11 m.grm.; HNO_3 , or 0.022 m.grm.; HNO_3 per cubic centimetre.

ACTION OF POTASSIUM PERMANGANATE UPON OXALIC ACID.

By ALBERT R. LEBDS, Ph.D.

In a paper entitled "Ozone and the Atmosphere,"† I stated that when the two solid substances, crystallised oxalic acid and potassium permanganate in powder, were mixed together, they liberated ozone on addition of water. Vacation intervening, the study of this and similar reactions was not resumed until five months later, in October of last year, when it was found that other organic acids yielded the same apparent results, more especially tartaric and acetic acids. The deportment of the inorganic acids varied: nitric acid, added to potassium permanganate, liberated a gas plentifully, which affected the potassium iodide papers, but the gas evidently contained nitrous acid, and when passed through water had no effect upon the test papers. Hydrochloric acid gave negative results, but chromic acid produced with the permanganate an apparently copious evolution of ozone.

An attempt was then made to determine the amount of ozone apparently given off. 20 grms. of pulverised oxalic acid were introduced into a half-litre flask, which was provided with a glass-funnel and stopcock for the introduction of a saturated solution of potassium permanganate, and an exit tube connecting with a wash-bottle con-

taining water. The corks used in this and the other experiments were boiled in paraffin, and the joints luted with the same. The escaping gas, after passing through the wash-bottle, was made to traverse a Geissler absorption apparatus, containing a decinormal solution of potassium iodide. The action was allowed to go on for a week at common temperatures, the permanganate being added in successive portions until the oxalic acid was completely decomposed. Nine litres of air were drawn through the apparatus, after which the flask was gently warmed and 9 litres more of air drawn through it. No change whatever had taken place in the potassium iodide. The entire experiment was carefully repeated with identical results.

This unexpected conclusion threw suspicion on the methods of ozone-generation, in which potassium permanganate is decomposed by an acid, and more particularly that in which sulphuric acid is employed. For this reason the same apparatus was used as in the above experiment, but 25 grms. pulverised permanganate were introduced into the flask and sulphuric acid run in through the funnel a drop at a time. Much caution was used in conducting the operation, but even then one flask was blown to pieces, after the reaction had been going on for several hours. In each of three trials the potassium iodide remained unchanged, but chlorine was found in the wash-water. This led to an examination of the reagents employed, with the result of finding that the potassium permanganate contained 0.164 per cent of potassium nitrate, and 7.47 per cent of other impurities, principally potassium chlorate, but along with it some manganic oxide, insoluble silicious residue, and a minute amount of potassium chloride. A trace of chlorine was likewise found in the oxalic acid. Analyses of other nominally chemically pure preparations were made with similar results, the least impure containing 97.06 per cent potassium permanganate, with 0.148 per cent chlorate, a trace of chloride, some manganic oxide and sand. The percentage of manganese, in the potassium permanganate first spoken of, determined gravimetrically by precipitation as sulphide, solution in hydrochloric acid, re-precipitation as carbonate and estimation as proto-sesquioxide, was 31.50 per cent, instead of 34.66, the theoretical amount. The manganese percentage determined volumetrically by means of ammonio-ferrous sulphate was 32.78, showing the influence of the admixture of chlorate upon the oxidation of the iron.

At this stage of the enquiry I found that the same sources of error had previously been exposed by C. Rammelsberg (*Berichte der Deutsch. Chem. Gesell.*, vi., p. 604), who showed that the potassium permanganate, which he experimented upon, contained only 27.3 per cent of manganese, the difference being due to 21.6 per cent of potassium perchlorate. He also found that when the permanganate, free from chlorine, was used, no smell was developed by the action of sulphuric acid, and the reaction upon iodo-starch was so weak that it evidently arose from a trace of chlorine.

ON THE WORKING OF MILD STEEL.

By SERGIUS KERN, M.E., St. Petersburg.

MANY experiments prove that mild steel, before being rolled into plates or bars, may be heated to a light welding heat, without the least fear of the ingot crumbling into pieces, which often happens with burnt steel. Altogether many persons are rather afraid of this method of working steel, and prefer to heat the ingots only to a white non-welding heat. This inferior mode of working has given rise to a large percentage of bad products, especially when plates are prepared, and plates with surface defects are certainly rejected. For cheap boiler and ship plates Bessemer steel is at the present time in constant use, and

* *Proceedings Amer. Chem. Soc.*, vol. ii.
† *CHEMICAL NEWS*, xxviii., p. 224.

the chief difficulty in obtaining clean plates in this case are the numerous blow-holes found in the outer parts of the ingots. The rolling of such ingots only masks these defects to a certain extent, as the small marks of their presence in the ingot may be always found in the finished plate. Often slag, scale, and small pieces of sand or brick laying on the surface of the ingot while it is in the furnace fall down into the blow-holes, and are rolled together with the ingot; such plates, with slag and brick particles rolled into the metal, must be certainly condemned for most purposes.

In order to avoid these inconveniences the following means are in the hands of the operator:—

1. The compressing of the liquid steel after casting in order to obtain sound ingots.
2. Hammering the ingots before the rolling of them.
3. Heating the ingots to a mild welding heat.

Compressing of the steel in liquid state must be effectual, but as the hydraulic press used for this purpose is very expensive, the ordinary prices of Bessemer steel, worked in such a manner, must be to a certain extent increased. Next the maintenance of the press, requiring a powerful engine and a couple of boilers, is also dear.

The hammering of the ingots before the rolling of them is cheap, and the work may be done in the Bessemer shop immediately after casting if a steam hammer may be placed in or near the casting house. The work requires a small amount of time, and the ingots, liberated from their moulds in 8 to 20 minutes after the casting of them, may be hammered, without placing them for this purpose into a furnace. After such an operation the surface blow-holes are smoothened and have no action on the surface of plates; this has been remarked during several trials.

The process of welding the steel is the cheapest way of doing away with the surface defects. The metal superficially welds, and when rolled gives a plate with a clean surface. For this purpose gas furnaces or ordinary reheating furnaces with blast may be used. The ingot is covered with sand and remains in the furnace till the scale runs off. Next it is rolled as quickly as possible. This method is even cheaper than the ordinary way of heating the ingots in furnaces without blast to a white non-welding heat. Certainly great care must be taken not to overheat the ingots, and a clever man must look after the furnace.

The burning of the carbon out of the metal is trifling during the heating of the ingot (1½ to 2 hours), as show the following analyses of plates worked by this method:—

Per cent of Carbon.

Ingots.	Plates.
0·24	0·20
0·24	0·19
0·25	0·21

Such plates have a beautiful shining surface, perfectly clean, and give a higher percentage of elongation.

A NEW FORM OF WASH-BOTTLE.

By M. H. FOYE.

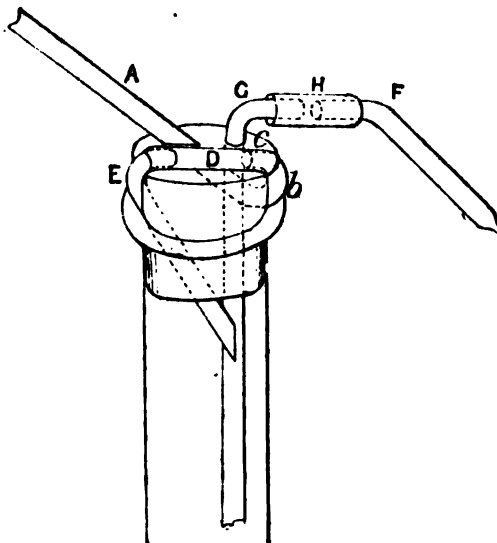
In this wash-bottle the blow-tube consists of three parts—A, D, E (see sketch). A and E are pieces of glass tube. A passes through the stopper obliquely, outside the bottle, is bent at *b*, and terminates at *c*. E is bent as shown, and passes through the stopper to the interior of the bottle. D is a piece of caoutchouc tubing which stretches across the top of the stopper and connects A and E. Now it will be seen that by the action of the forefinger upon D, the blow-tube can be closed and opened at pleasure. The advantages of this arrangement are the following:—

1. That after you create a pressure on the liquid in the bottle by flowing through the blow-tube, you can, by compressing the caoutchouc tubing, maintain

this pressure, without continual blowing, until it expends itself by expelling liquid from the jet.

2. The disagreeable reflux of steam into the mouth when washing with hot water, of ammonia or acid fumes when using solutions of either of the latter, can be completely prevented.
3. The flow of liquid from the tube A, when it is desired to use the bottle in that way, can be so accurately regulated by pressing the caoutchouc tubing that it is particularly valuable for diluting, filling measuring vessels, &c.

The jet I use, and which has been suggested by a friend, needs a word of explanation. It is bent, as shown, at *F*, and attached to the tube G, which dips into the liquid in the bottle, by a thick piece (thick enough to maintain



the jet in position) of caoutchouc tubing, H, by turning which round G, the jet can be directed upwards, downwards, or sideways, as desired.

This wash-bottle can be used in every respect like the ordinary wash-bottle, and as conveniently, besides having the advantages described above. It has been used for the last nine or ten months by myself and others in the laboratory at Crewe, and has given every satisfaction. It is not at all liable to get out of order, and is but slightly more difficult to make than the ordinary form of bottle.

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REACTIONS OF
IODINE AND OF POTASSIUM IODIDE
WITH SULPHUROUS ACID.

By ALBERT E. MENKE.

JOHNSON, in his investigation of potassium tri-iodide, noticed that on mixing strong solutions of potassium iodide and sulphurous acid, a yellow colour was produced, which was not bleached by more sulphurous acid, but disappeared on adding water. Gmelin ("Handbook of Chemistry," Watts, ii., 263) states, on the authority of Saladin (*Journ. Chim. Med.*, vii., 528), that hydriodic and sulphurous acids in aqueous solution form a yellow liquid, brighter in proportion to the concentration, and from which, eventually, sulphur is separated. Having myself verified this statement, I inferred that the yellow colour

given by potassium iodide with sulphurous acid was due to the action of the latter upon the hydriodic acid set free, and I was anxious to isolate the substance causing the colour. That it could not be iodine was shown by its resisting the action of excess of sulphurous acid, and by its not colouring carbon disulphide. When the yellow solution was shaken with ether, the latter was very slightly coloured, but on adding a little alcohol and again shaking the ethereal layer acquired a bright yellow colour, and when drawn off and evaporated it left a brown oily substance, having a pungent odour somewhat resembling that of bromine. On heating moderately it evolved vapour of iodine. I made several attempts to obtain it in a state fit for analysis, but the yield was always small, and there was a tendency to the separation of plates of iodine. I therefore tried to obtain it by the action of sulphurous acid gas upon iodine dissolved in ordinary alcohol, hoping that, the quantity of water present being very small, a larger quantity of the yellow substance would be produced. One ounce of iodine was dissolved in 20 c.c. of alcohol, and sulphurous acid was passed in to complete saturation; the liquid remained of a dark brown colour, which disappeared immediately on adding excess of water. As it did not deposit anything after some days, and was found to contain a little free iodine, small quantities of water were added, and more sulphurous acid gas was passed in until no more free iodine could be detected, although the liquid had still a dark brown colour. On standing it deposited a plastic substance resembling the amorphous form of sulphur. This was freed by pressure from adhering mother-liquor, when it weighed 0.140 grm. On oxidising it with fuming nitric acid and precipitating with barium chloride it gave barium sulphate corresponding to 60 per cent of sulphur. Though disappointed in isolating the substance, I submit that my experiments justify the conclusion that the yellow body formed when sulphurous acid acts upon concentrated solutions of potassium and sodium iodide is an unstable iodide of sulphur, and I would suggest the following equations explaining the reaction:—

- (1.) $KI + H_2SO_3 = HI + KHSO_3$.
- (2.) $8HI + 2SO_2 = 4H_2O + I_6 + I_2S_2 (?)$
- (3.) $I_2 + 2H_2O + SO_2 = 2HI + H_2SO_4$.

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NOTICES OF BOOKS.

A Treatise on Chemistry. By H. E. ROSCOE, F.R.S., and C. SCHORLEMMER, F.R.S. Vol. II. Metals. Part I. London: Macmillan and Co.

THIS is one of those books most valuable to the student, but the despair of the reviewer. We have no novel and questionable theories to criticise, no new facts of capital importance to point out. On the other hand, it will not be expected that the work of two chemists of such eminence can be rich in errors. We might, indeed, on careful examination come upon statements which we should hesitate to endorse. We may here and there think that some piece of information which has been omitted ought to have been given, and it would certainly be no difficult matter to select from the pages before us interesting facts not to be met with in those chemical hand-books, manuals, and treatises which have lately become somewhat too numerous. The book has, indeed, a distinctive character. Its illustrations, in their number, excellence, and accuracy, have few rivals. They are, further, not restricted to sketches of the apparatus used in the scientific laboratory or at the lecture-table, but include careful representations of the furnaces and other plant as now actually used in the best-arranged chemical manufactories and metallurgical establishments, with the additional recommendation of being drawn to scale. Such impor-

tant processes as the manufactures of alkali and of bleaching-powder, are also described, if not so as to enter into every nicety, still in a manner which has in it no trace of the careless and the perfunctory. In these important respects the work is advantageously distinguished even as compared with such valuable treatises as those of Gmelin and Watt. Let any one, for instance, read the brief section on artificial soda in Gmelin (vol. iii., p. 79), and he will know how to appreciate the difference. Even some professedly technological works give less thorough descriptions of the various stages of the alkali manufacture than does the work before us. This feature is, we hold, of extreme value. Before the scientific chemist can bring his theoretical knowledge to bear usefully upon any of the industrial arts he should have an accurate general knowledge of their nature; of the various portions of the plant and of their functions, and of the difficulties which have been, or which still have to be overcome. Now, without wishing to convey the impression that mere reading can at once put the student in possession of practical knowledge, we hold that a young man fresh from college and entering upon the position of a "work's chemist," will find himself much sooner at home and do himself more credit if the work of Profs. Roscoe and Schorlemmer has been his guide than if he has merely been accustomed to manuals which scarcely condescend to refer to manufacturing operations. We believe that "theory with practice" is the key-word to industrial success, and in that conviction we congratulate the authors on their undertaking.

Coal, its History and Uses. By Professors GREEN, MIALI, THORPE, RUCKER, and MARSHALL, of the Yorkshire College. Edited by Prof. THORPE. London: Macmillan and Co. 1878.

TOWARDS the close of the year 1877 it was suggested to the Professors of the Yorkshire College that by delivering a series of lectures in some of the larger towns of the West Riding they might aid in the good work which is being done by Dr. Gilchrist's educational bequest. Each lecturer was fain to choose his own topic, the whole of them agreeing that, considering the locality in which they were to lecture, they could not select a better general subject than coal. Early in 1878, therefore, a series of ten lectures was delivered in Leeds and Keighley in the following order:—Lectures I. and II., on the Geology of Coal, by Prof. Green; III. and IV., on Coal Plants and Animals, by Prof. Miall; V. and VI., on the Chemistry of Coal, by Prof. Thorpe; VII. and VIII., on Coal as a Source of Warmth and Power, by Prof. Rücker; and IX. and X., by Prof. Marshall, on the Coal Question. The present volume consists of these ten lectures altered in form, revised and arranged consecutively in ten chapters. The only ones affecting our readers are those of Prof. Thorpe, who has managed to introduce a certain amount of fresh detail into this most hackneyed subject by giving a popular account of Mr. J. W. Thomas's researches on the gases occluded in coal. He also gives an interesting description of the successive steps in the invention of the Davy lamp, illustrated by diagrams. Mr. Ansell's fire-damp alarm is also fully described and illustrated. The lectures are illustrated with nearly sixty well-executed cuts. A full index is appended.

Year-Book of Pharmacy. Comprising Abstracts of Papers relating to Pharmacy, Materia Medica, and Chemistry, contributed to British and Foreign Journals from July 1, 1877, to June 30, 1878, with the Transactions of the British Pharmaceutical Conference at the 15th Annual Meeting. London: J. and A. Churchill, 1878.

THE ninth volume of the "Year-Book of Pharmacy," as usual, has appeared somewhat late. The book contains abstracts of papers relating to pharmacy contributed to British and foreign journals for the year ending June 30,

1878, and the Proceedings of the British Pharmaceutical Conference which held their sittings in the middle of August. According to the statement which Prof. Attfield made at the meeting held on August 12th, the MS. of the book was to have been laid upon the table on the following day; we cannot, therefore, understand why so long a period as three months and a half should be allowed to elapse before the appearance of the book, even supposing that the Transactions of the Conference were not included in the MS. mentioned by Prof. Attfield. The first part of the work is, as usual, particularly copious and complete, and gives us all the information contained in the principal pharmaceutical and medical journals issued in England and America during the period referred to in a condensed form.

The second part of the work, containing the Proceedings of the Conference, shows that the working members of that body have applied the wholesome lesson read to them by Prof. Attfield at the 1877 meeting, and we are glad to see that every paper, without exception, is of a thoroughly practical character from the pharmacist's point of view. The place of honour is occupied by the paper of Dr. Alder Wright and Mr. A. P. Luff, on the aconite alkaloids. Those improved processes for purifying the aconite bases have led to unexpected results. Thus, the substance described by them last year under the name of pseudaconitine is not a definite base, but a mixture of pseudaconitine and apopseudaconitine, the latter being a dehydrated derivative of the former. The substance again which was last year regarded as pseudaconine is now distinguished as apopseudaconine, and is a dehydrated derivative of true pseudaconine. True aconitine also forms a similar series of dehydrated derivatives. The veratrum alkaloids seem likely to afford a similar series of bases. The alkaloid or alkaloids obtained from *aconitum ferox* are still a puzzle. In any case they appear to differ both from aconitine and pseudaconitine. The outcome of all these apparently recondit researches is of the highest possible practical importance, hitherto the pharmacist has been unable to present the physician with any preparation of either British or foreign aconite root which had a constant composition, but now, as the result of the investigations of Dr. Wright and his colleagues, the active principle of the drug will be procurable in a form whose composition will be nearly as definite as that of potassium nitrate. Mr. Shenstone's researches on the action of nitric acid on strychnine seem to disprove Prof. Sonnenschein's alleged discovery of the formation of brucine by this method, and would also lead us to suppose that pure brucine has not yet been procured. Most of the other papers are of too technical a nature for notice in our pages.

A Pocket-book for Chemists, Chemical Manufacturers, Metallurgists, Dyers, Distillers, &c. By THOMAS BAYLEY, Assoc. R.C.Sc.I., Demonstrator of Practical Chemistry, &c., in the Mining School, Bristol. London: Spon and Co. 1878.

A HANDY little book that has long been wanted, and of whose usefulness it would be needless to say anything. Hitherto chemists have been obliged to fall back on the very incomplete little note-book of Gutch or the French *Addenda* of Dunod, and it seems surprising that the want has never been supplied. The tables, of course, begin with a list of the elements, their symbols, atomicities, and atomic weights; but why, we may ask, are the latter repeated twice? once according to the latest determinations, and again according to some obsolete table. The space thus wasted might have been devoted to the date of the discovery of the newer elements and the names of their discoverers. Various other tables, data, and formulæ follow, one of the most useful being a table of coefficients giving the amount of the constituent sought by simple multiplication. Another is a table for the conversion of grammes into grains and *vice versa*, from 1000 grammes to

1 milligramme, and from 1 grain to 1000 grains. The thermometrical tables range from 263° C., 208° R., and 500° F., down to -10° C., -1° R., and +14° F. The barometrical tables range from 27 inches to 30.98 inches. Boiling points, specific gravities, vapour densities, and solubilities follow. The tables showing the specific gravities of solutions of salts, &c., of different strengths, are particularly copious, but we cannot understand why it is that the temperature at which the specific gravities have been taken differ in every instance. The analytical tables are also very full. The pages on chemical manipulation might, we think, have been omitted. The glossary of minerals, giving their names, formulæ, hardness, specific gravity, and crystalline system, will render Mr. Bayley's little book indispensable to the mineralogist. We have throughout tested Mr. Bayley's *vade mecum* for tables of every sort and kind, and have every one down even to a ready-reckoner and wages table. The 421 pages which the book contains literally team with information, and we think it will puzzle Mr. Bailey to add very much to his second edition, which cannot fail to appear very speedily. The type in which the tables are printed is clear, though necessarily small in cases where much matter is crowded together, and there appear to be but few misprints.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 24, December 9, 1878.

Artificial Pyroxene (Diopside).—L. Gruner.—Crystals of this mineral have been obtained from bricks very rich in lime and magnesia, after being exposed for some time to an intense heat in a furnace.

Influence of Atmospheric Electricity upon the Fructification of Plants.—L. Grandeau.—Atmospheric electricity in two sets of experiments conducted at Nancy and at Mettray decidedly promotes fructification.

Disease of the Coffee-tree Observed in Brazil.—C. Jobert.—This affection, the symptoms of which are minutely described, attacks chiefly plantations in moist and shady situations, and appears to be caused by *Anguillula* which attack the roots.

Diffusion of Heat by Leaves.—M. Maquenne.—The green organs of plants diffuse a considerable proportion of the heat-rays which they receive, this action being almost always accompanied by imperfect reflection, the reflected rays being polarised in the plane of incidence. The proportion of rays diffused in the case of normal incidence diminishes with the temperature of the source of heat.

The Power of Kinds of Wood in Absorbing Water.—E. J. Maumené.—The author finds the proportion of water absorbed by different sorts of wood varies from 9.37 to 174.86 per cent.

No. 25, December 16, 1878.

Observations on M. Pasteur's Paper on Alcoholic Fermentation.—M. Berthelot.—The author, feeling himself reflected upon by M. Pasteur's remarks on the posthumous essays of Claude Bernard, has undertaken a novel and interesting experiment. He writes, "when speaking of a soluble alcoholic ferment capable of being consumed step by step with its production and in the very chemical act which it determines, I took care to add that, for the demonstration of this hypothesis, it would be necessary to discover the conditions in which this ferment is produced in more considerable proportions than the quantity destroyed in fermentation. Such were the con-

ditions which C. Bernard seems to have met with in the experiments, the account of which has unfortunately reached us in so imperfect a manner. Nevertheless, I considered it useful to science to publish them, such as they are, with the object not of opening a controversy but of pointing out a new path for research." By simultaneously hydrogenising and oxidising sugar M. Berthelot has in fact succeeded in producing alcohol, though in very small proportions. He considers that his present results do not warrant a definite conclusion, for the limit may be due as much to the inaccuracy of the fundamental hypothesis as to the imperfection of the conditions. Still, he considers the fact that alcohol has been produced in the cold by the electrolysis of a solution of sugar merits to be placed on record.

Novel Phenomenon of Static Electricity.—E. Duter.—In reply to the memoir of M. Govi (December 2) the author does not admit that his researches can be viewed as a continuation of those of the Italian physicist.

Artificial Production of Nepheline and Amphigene.—F. Fouqué and A. Michel Lévy.—These minerals have been obtained by the same method used by the authors for the reproduction of feldspars.

Spectrometric Measurement of High Temperatures.—A. Crova.—The spectrometric study of the luminous radiations emitted by incandescent bodies has led the author to the discovery of a new method of determining elevated temperatures by the analysis of the light which they emit. If we take, in the continuous spectra of light emitted by two incandescent sources, the one of known temperature, T , and the other of unknown temperature, x , two simple radiations of very different wavelengths, λ and λ' , to which we may refer all our measurements, and determine by means of a spectro-photometer the ratios—

$$\frac{I}{I'} \text{ and } \frac{i}{i'}$$

of the intensity of the two rays λ and λ' in the two spectra. The quotient of these two ratios represents the ratio of the intensities of the ray λ' in the two spectra when the more intense has been lowered so as to give the same intensity to the ray λ in the two spectra.

Specific Heat and Melting Temperature of Palladium.—J. Violle.—The mean specific heat of palladium at $100^\circ = 0.0592$. Its melting temperature is 1500° .

Influence of Temperature on Rotatory Magnetic Power.—J. Joubert.—In the author's experiments the rotatory power was found to increase with the rise of temperature.

Density of Coefficients of Expansion of Liquid Chloride of Methyl.—MM. C. Vincent and Delachanal.—The sp. gr. at $-23.7^\circ = 0.9945$; at $+39^\circ = 0.87886$. The values found for the three coefficients α , β , γ , respectively, are 0.00193929 , 0.0000183121 , and 0.00000105916 .

Oxidation of Certain Aromatic Derivatives.—A. Etard.—On oxidising various organic compounds by means of chromic chlorhydrate the author observed very different reactions, and found that its mode of action is principally regulated by the nature of the body acted upon and by that of the substituted groups which it contains. Thus there are formed acetons, aldehyds, and quinons, whilst hydrochloric acid, or even chlorine, is liberated as an accessory product; in other cases there is no escape of gas, an immediate and total combination being formed.

Nature of Certain Crystalline Secondary Products Obtained in the Industrial Treatment of Pennsylvanian Petroleum.—L. Prunier and R. David.—Among these accessory products are found the carburetted derivatives of acetylen and benzin (anthracen, chrysen, &c.) discovered in the products of the distillation of coal.

Hæmocyanin, a New Substance from the Blood of Octopus vulgaris.—L. Fredericq.—The liquid portion of the blood of the cuttles contains a colourless albumenoid substance, forming with oxygen an unstable compound of a deep blue colour. This substance plays, in the respiration of the cuttles, the same part as does hæmoglobin in that of the Vertebrata. The veinous blood of the cuttles is colourless, but the arterial is a dark blue. If hæmocyanin is ignited it leaves an ash rich in copper, which seems to be in the same condition and to fulfil the same functions as does iron in hæmoglobin.

Influence of the Different Colours of the Spectrum on the Development of Animals.—E. Yung.—Referring to earlier researches on this subject the author states that M. Beclard placed the eggs of *Musca carnaria* under glasses of different colours, and remarked that they were developed very unequally, those under the blue and violet ray being the most developed, and those under the green ray the least so. He arranges the rays, as regards the development of larvæ, in the following order:—Violet, blue, red, yellow, white, green. The author has carried on for three years a series of observations on the eggs of the common and esculent frog, of the trout, and of *Lymnaea stagnalis*. These eggs were placed in vessels plunged respectively into violet, blue, green, yellow, red, and white solutions, whilst one vessel was kept in a dark closet. Violet light accelerates the development in a remarkable manner, and is followed in this respect by the blue, the yellow, and the white. Red and green light appear injurious, as the author was not able to obtain the complete development of ova in these colours. Darkness did not hinder development, but retarded it, contrary to the results of Higginbottom and MacDonnell. The colours may be arranged in the following series of diminishing activity:—Violet, blue, yellow, and white (which are nearly equal), darkness, red, and green. Tadpoles deprived of food died more rapidly of inanition under the violet and blue rays than under the others. The general mortality seemed lowest in white light.

Gazzetta Chimica Italiana.
Anno viii., 1878. Fasc. vi. and vii.

Crystalline Form of Usnic Acid.—P. Freda.—The same remark applies to this memoir.

Preparation of Digallic Acid.—Dr. P. Freda.—A criticism on the method proposed by Prof. Hugo Schiff. The author finds that on heating gallic acid with arsenic acid, whether in alcoholic or aqueous solution, a substance is obtained having some reactions similar to tannic acid. This substance constantly contains arsenic acid, and if this is removed the compound is re-converted into gallic acid.

On α -Iso-chloro-butyric Acid and some of its Derivatives.—L. Balbiano.—The derivatives are oxy-iso-butyric, metacrylic, and dibutylic acids.

New Phenomena Observed in "Plastering" Wines and Musts.—Prof. E. Pollacci.—Sulphate of lime reacts solely upon the cream of tartar, producing acid sulphate of potassa, which remains in solution, while tartrate of lime is formed and is chiefly deposited. The reaction between the two salts is not complete, since there may be found in the liquid both sulphate of lime and cream of tartar. If the gypsum is in large excess a part of it is deposited among the tartrate of lime.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 14, December 5, 1878.

Agoraphobia.—Dr. Legrand du Saule.—The author has studied a novel nervous affection, which manifests itself in a dread of open spaces.

Biedermann's Central-blatt.
Heft 12.

Researches on the Influence of Artificial Manures upon the Growth of Summer Barley.—Dr. G. Marek.—The author concludes that on fruitful soils artificial manures are untrustworthy for barley. Unmanured plots often yield better results than such as are manured.

Experiments with Sugar Beets in Different Soils and with Different Manures.—Dr. J. Hanamann.—The author draws no conclusions from his experiments.

Composition of Wool and Hair.—P. Schutzenberger.—From *Comptes Rendus*, 1878, pp. 767-69.

On Certain Animated Enemies of Cultivated Plants and their Destruction.—M. Schaefer.—An account of certain insects hostile chiefly to the vine. Among the remedies proposed are sodium sulphide, petroleum, and a mixture of an alcoholic extract of tobacco, with bisulphide of carbon.

Wine-Testing and Chemical Analysis.—Drs. E. Mach and Patele.—The authors show that wines on analysis often display properties not suspected from their taste. Sorts rich in alcohol are often pronounced light, and others strongly acid seem mild, &c. These seeming contradictions are due to the fact that wine is a most complicated liquid, whose components are by no means all known. Even among the recognised and more prominent ingredients, it is not so much the quantity of one or the other as their relative proportion which determines the flavour.

On the Influence of Temperature in Malting.—A. Prantl and Hans von der Planitz.—Not susceptible of useful abstraction.

Oino-Chemical Researches.—Dr. M. Buchner.—In order to solve the question whether a given wine is natural or fictitious, in the former certain proportions prevail between the quantities of extract, alcohol, and ash. If the relative proportion of these constituents vary from the normal standard, or if the total is too high or too low, the case is suspicious.

A Source of Error in the Determination of Fat in Milk.—Dr. L. Maneth and Dr. G. Musso.—The authors maintain that if milk, evaporated down along with sand, is extracted with ether, the ether driven off, the residue again extracted with ether and driven off, and the final residue extracted with sulphide of carbon, there remain certain strongly acid drops, of a yellow or deep red colour, soluble in ether and water, but insoluble in fat or in sulphide of carbon. At 100° they take a darker colour and become almost resinous. The authors consider them as impure lactic acid. In recent milk there is little of this substance, and hence there is very little difference between the weight of the residue from the original extraction with ether and that of the residue from the evaporation of sulphide of carbon. There is more difference in old milks, and especially in cheese. The authors state that in milk evaporated to dryness at high temperatures there occurs a wax-like matter, sparingly soluble in ether, which greatly impedes and protracts the process of extraction. (The editor of the *Milch-Zeitung*, from which the above paper is taken, does not find a double extraction of the milk residue with ether requisite. If a proper apparatus is used, the first ethereal extract is perfectly clear, and leaves on evaporation clear fat, in which no drops of foreign matter are present).

Investigations on the Nitrogenous Food of Fungi.—Th. Schlösing and A. Müntz.—From the *Comptes Rendus*, 1878, p. 892.

Analysis of Stag and Roe Horns and of the Horns of an Ox.—Prof. W. Vesely.—Stag, 50.480 ash, 7.88 moisture; fallow-deer, 50.640 ash, 8.98 moisture; roe, 47.830 ash, 9.42 moisture; ox, 1.167, 8.75 moisture—the difference in each case being organic matter. The ash contained:—

	Stag.	Fallow deer.	Roe.	Ox.
Lime	51.96	52.29	52.13	31.60
Magnesia ..	1.02	0.96	1.26	3.60
Phosph. acid ..	42.19	41.41	40.60	12.60
Carbonic acid ..	2.12	2.29	2.98	—
Potash	0.56	1.13	0.89	5.50
Soda	0.30	—	—	—
Sulph. acid ..	0.18	0.09	0.07	6.80
Fluorine	1.56	1.48	1.96	—
Silica	0.04	0.05	0.09	1.90
Ferric oxide ..	0.79	0.30	0.02	20.30 (?)
Carbon	—	—	—	1.36

MISCELLANEOUS.

Russian Scientific News.—At the Technical Society of St. Petersburg M. Latchinoff delivered a very interesting lecture on the electric light. He made some experiments with Jablochkoff's condensators, which consist of a set of tin plates placed one on another; the surface of every plate is 0.7 square metre. Between every pair of such plates a piece of silk covered with varnish is introduced. The height of the condensator was about 6 feet. On introducing two condensators into a circuit the intensity of the electric light is doubled. Such condensators are not cheap owing to the great quantity of silk wanted, and thus the application of this apparatus is limited. The lecturer believes the new system of electric lighting devised by M. Rapiéff to be a serious opponent of Jablochkoff's process. The chief advantage of the new system is that the luminating point does not change its position, and therefore this system is more suitable for the projection of the electric light at a distance. This advantage will give increase to the use of the electric light for military purposes.

M. Gregorieff, of the Petrovsky Agricultural Academy of Moscow, has made some interesting experiments concerning the quality of milk sold in various places in the city of Moscow. In large towns of Europe the falsification of milk has given rise to a great number of investigations, and this question was discussed in detail. It appears that in Moscow the milk sellers mix water very moderately in a few cases, and have altogether little knowledge about the "chemicals," viz., chalk, starch, flour, &c. In all the samples analysed, no such "chemicals" were present. The 64 samples of milk were taken from the following places in Moscow:—Samples A, from peasants living in the vicinity of the town (17); samples B, milk from the town cows (6); samples C, from small grocers' shops (14); samples D, from milk shops (22); samples E, milk of the Academical farm.

The following are the average results of the analyses of these milks:—

	A.	B.	C. Samples.	D.	E.
Water	88.13	88.08	89.16	87.48	87.53
Dry substance..	11.87	11.92	10.84	12.52	12.47
Fat	2.76	2.39	1.55	3.13	4.10
Mineral substance ..	0.87	0.79	0.71	0.69	0.67

As compared with an average analysis of normal milks from healthy cows, the first four samples cannot be called bad specimens. For comparison the following analysis (average) may be mentioned:—

Water	87.25
Dry substance ..	12.75
Fat	3.50
Mineral substance ..	0.75

These results were obtained by Fleischmann (Molkereiwesen, 1876). The milk of the Academical Farm (sample E) is certainly a very high and good specimen; such milk will never be found in the hands of the sellers.

S. KERN, M.E.

St. Petersburg.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

DECEMBER, 1878.

THE following are the returns of the Society of Medical Officers of Health:—

Appearance in a foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	Sulphuric An- hydride.	Hardness on Clark's Scale.	
	Saline.	Organic.								Before Boiling.	After Boiling.
	Grs.	Grs.								Grs.	Degs.
<i>Thames Water Companies.</i>											
Grand Junction Clear	0'000	0'010	0'180	0'075	23'70	8'400	0'684	0'64	1'240	15'9	4'00
West Middlesex Clear	0'000	0'010	0'090	0'089	23'30	7'800	0'648	0'72	1'860	14'3	4'20
Southwark and Vauxhall Clear	0'000	0'006	0'120	0'115	22'20	7'110	0'720	0'72	1'400	14'3	4'60
Chelsea Clear	0'000	0'010	0'120	0'073	22'50	6'720	0'720	0'72	1'600	14'8	5'10
Lambeth Clear	0'000	0'010	0'105	0'103	22'00	7'040	0'576	0'72	1'630	14'3	5'10
<i>Other Companies.</i>											
Kent Clear	0'000	0'002	0'360	0'002	29'50	11'560	1'081	1'152	3'300	17'6	5'10
New River Clear	0'000	0'005	0'165	0'057	21'40	7'400	0'576	0'664	1'060	14'8	4'20
East London Clear	0'000	0'009	0'105	0'065	23'00	7'800	0'720	0'864	1'160	14'8	5'10

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

MEETINGS FOR THE WEEK.

- MONDAY, Jan. 13th.—Medical, 8.30.
London Institution, 5.
Geological, 8.
- TUESDAY, 14th.—Civil Engineers, 8.
Photographic, 8.
Zoological, 8.30.
Royal Institution, 3. "Animal Development,"
Prof. Schäfer.
- WEDNESDAY, 15th.—Meteorological, 7.
Society of Arts, 8. "Economy and Safety by
the Use of Automatic Couplings on Rail-
ways," T. A. Brocklebank.
- THURSDAY, 16th.—Royal, 8.30.
Chemical, 8.
Royal Society Club, 6.30.
Royal Institution, 3. "Electric Induction," J. H.
Gordon.
- FRIDAY, 17th.—Royal Institution, 9. "Electric Light," Prof. Tyndall.
Society of Arts, 8. "Afghanistan," C. E. D. Black.
- SATURDAY, 18th.—Royal Institution, 3. "Reptilian Life," Prof. H.
G. Seeley.

INSTITUTE OF CHEMISTRY OF GREAT
BRITAIN AND IRELAND.

An Examination in Practical Chemistry in connection with the Institute of Chemistry will be held during the last week of January next. Examiner, Dr. W. J. Russell, F.R.S. Candidates can obtain further information on application to the Secretary, Mr. Charles E. Groves, Somerset House Terrace, London W.C.

The Authors of "A Practical Treatise on the
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TO IRONMASTERS AND OTHERS.

An Associate of the Royal School of Mines, who has been Chemist for the last eight and a half years at Iron Works in Northamptonshire, wishes for a re-engagement as Manager or Chemist. Is well acquainted with the working, analysis, and smelting of Northamptonshire ironstone, and has been accustomed to teach analysis and assaying to pupils in the laboratory.—Address, "Chemist," Midland Road, Wellingborough.

Situation wanted by a Competent Analyst as Assistant or Principal. Good teacher. Apply, Public Analyst, CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

Wanted, a Foreman in Superphosphate Works. Capable of undertaking the making of Sulphuric Acid and mixing. First class references required. He must be a married man. Apply to Edward Toyne, Saxilby, Lincoln.

THE
QUARTERLY JOURNAL OF SCIENCE.

Edited by WILLIAM CROOKES, F.R.S., &c.

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- I. On the Thickness of the Antarctic Ice, and its Relations to that of the Glacial Epoch. By James Croll, LL.D., F.R.S.
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NOTICE.

MONTHLY ISSUE OF
THE JOURNAL OF SCIENCE.

The JOURNAL OF SCIENCE will in future be issued MONTHLY instead of QUARTERLY, and will consist of 48 pp., the form and general appearance remaining the same.

The first of the Monthly issue will appear on
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THE CHEMICAL NEWS.

Vol. XXXIX. No. 999.

ESTIMATION OF CYANOGEN IN SODA-LYES.

By FRED. HURTER Ph.D., &c.

THE cyanogen compounds which occur in soda-lyes are the following:—

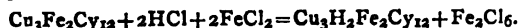
Sodium ferrocyanide,
Sodium sulphocyanide,
Sodium cyanate.

Though in crude soda (black-ash) the cyanogen is contained as cyanides, the soda-lye obtained therefrom contains no cyanides. During the lixiviation the cyanides are transformed into ferrocyanides, and all the vat-liquors I have yet investigated contained more iron in solution than the cyanogen would account for. If, however, sodium cyanide is present at all it is readily converted into ferrocyanide by boiling the liquid with freshly precipitated protoxide of iron.

The most important of the cyanogen compounds to the manufacturer is the ferrocyanide, since this salt brings into the solution a sufficient amount of iron to colour the finished product slightly brown.

The sulphocyanide and the cyanate are practically of no importance, as they yield colourless products of decomposition during the further treatment of the soda. The chief problem to the chemists of alkali works, consists, therefore, in the rapid estimation of the ferrocyanides. The following method will be found exceedingly rapid, and sufficiently accurate for solutions containing not more than a grms. of sodium cyanide per litre.

When soluble ferricyanides are mixed with salts of copper a yellow precipitate of ferricyanide of copper is formed. If a proto-salt of iron is added afterwards a blue precipitate will be formed so long as any ferricyanide exists still in solution. As soon, however, as all the ferricyanide has been precipitated, the addition of a ferrous salt produces no longer a blue colour, but reacts upon the ferricyanide of copper, reducing it to ferrocyanide of similar composition to that obtained on adding to an excess of ferrocyanide of potassium a copper solution gradually. The reaction is represented by the following equation:—



This reaction suggests an indirect estimation of the cyanogen compounds by measuring the amount of protoxide of iron which can be transformed into peroxide. It is not, however, this which forms the principle of the method now to be described, and which makes use of the above reaction simply as indicator.

100 c.c. of the strong soda-lye are oxidised by means of chlorine, hypochlorite of soda, or, simpler, bleaching-powder solution, until the whole of the sulphides, &c., are converted into sulphates, and the ferrocyanide into ferricyanide; the solution, after being acidified and freed as much as possible from excess of chlorine by warming and agitating, is ready for titrating.

On a porcelain slab sprinkle a few drops of a dilute solution of ferrous sulphate (1 part of ferrous sulphate to 100 parts of water). Add now to the solution to be analysed a twentieth normal copper solution from a burette until a drop of the solution, on being brought in contact with a drop of ferrous sulphate solution, no longer gives a blue colour, but yields the pure purple colour of cupric ferrocyanide. The copper solution is prepared by dissolving pure metallic copper in as little nitric acid as possible, and diluting with distilled water. (3.17 grms. of

copper are diluted to 1 litre. 1 c.c. = 0.001313 grm. Na_4FeCy_6 .)

This process, from its extreme simplicity, will commend itself for approximate comparative examinations.

The ferrocyanide of copper, if present in larger quantities than those indicated above, hides too much the blue colour of the Prussian blue, and must be separated before applying the ferrous sulphate. Moreover, in concentrated solutions, the precipitate of ferricyanide of copper contains more cyanogen per atom of copper than the formula $\text{Cu}_2\text{Fe}_2\text{Cy}_{12}$ requires. In dilute solutions, however, the amount of copper solution required is such as to correspond sufficiently close with the formula above stated; and considering that not one of the many other compounds contained in soda-lyes interferes with this process of estimating the cyanogen, it will rival in accuracy any other volumetric process devised for this purpose.

If it is apprehended that the solution under examination contains cyanide of sodium beside ferrocyanides, it must first be boiled with a small quantity of ferrous oxide, produced by adding a little ferrous sulphate. The solution is then oxidised, acidified, and is ready for testing.

The sulphocyanide is present in traces only. It can approximately be estimated by acidifying the solution under examination, adding some chloride of zinc to precipitate ferrocyanide of zinc. The solution is then filtered, and the filtrate coloured by means of ferric chloride. In a second vessel an equal quantity of ferric chloride, diluted to the bulk of the solution under examination, is coloured by means of a solution of sulphocyanide of potassium of known strength, until the tint of both solutions is alike. The amount of sulphocyanide thus consumed is in some degree a measure of the sulphocyanide contained in the lye, but this is usually so small that its estimation is of no practical utility.

Laboratory of Messrs. Gaskell, Deacon, and Co.,
Widnes, January, 1879.

ON THE COMBINATIONS OF AURIN WITH MINERAL ACIDS.*

By R. S. DALE, B.A., and C. SCHORLEMMER, F.R.S.

In our last communication† we stated that by the action of acetyl chloride on aurin we obtained a colourless crystalline compound, which we intended to examine more closely. We have since found that this body is identical with a compound which Gräbe and Caro‡ obtained by the direct union of aurin and acetic anhydride and having the formula $\text{C}_{19}\text{H}_{14}\text{O}_3 + \text{C}_4\text{H}_6\text{O}_3$.

We also mentioned that the purification of this substance was found to be beset with several difficulties. The cause of this was found out after some trouble, but at the same time we were rewarded by the discovery of a series of remarkable bodies, consisting of combinations of aurin with mineral acids.

These salts, as we may call them, are beautiful bodies, crystallising exceedingly well, and although some of them are decomposed by water, they are very stable in dry air. To their discovery we were led by the following observations.

On heating aurin with glacial acetic acid and acetyl chloride, the crystals lose at once their steel-blue lustre and assume a pale red colour. To obtain the compound thus formed in a pure state, acetyl chloride was added to a saturated solution of aurin in acetic acid. The liquid assumed at once a much lighter colour, and soon pale red needle-shaped crystals having a diamond lustre separated out. On re-crystallising these repeatedly from alcohol

* Read before the Manchester Literary and Philosophical Society December 20, 1878.

† *Proc. Lit. and Phil. Soc.*, 1878, 141, and *CHEM. NEWS*, vol. xxxviii., p. 34.
‡ *Ber. Deutsch. Chem. Gesell.*, xl., 1, 122.

we obtained oblong six-sided plates, which, as analysis showed, were pure aurin.

On treating the original crystals with water they become dull and brownish red, the solution containing acetic and hydrochloric acids. It therefore seemed not improbable that an additive product of aurin and acetyl chloride had been formed, containing, however, also acetic acid, as a superficial examination showed that the liquid contained, to one molecule of hydrochloric acid, much more than one molecule of acetic acid. We therefore tried to obtain an analogous benzoyl compound, and to determine in it, after decomposition with water, the relative quantities of hydrochloric and benzoic acids.

On adding benzoyl chloride to a hot solution of aurin in acetic acid, similar crystals as before were obtained, which, after being dried on filter-paper in dry air, were decomposed by water, but only hydrochloric and acetic acid went into solution, and on heating the product with water or alkalis but a mere trace of benzoic acid could be detected.

These facts, coupled with the observation that the bright red needles which, as we stated in our former paper, are formed by crystallising aurin from hot aqueous hydrochloric acid, retain the latter obstinately, led us to the conclusion that this acid forms a definite compound with aurin.

Such a body could be formed under the above conditions, as our glacial acetic acid contained a little water. Moreover, Mr. Charles Lowe had informed us that the splendid specimen of aurin which he exhibited at Paris was obtained in the following way:—The crude but crystalline aurin, which is obtained by heating pure phenol with sulphuric and oxalic acids, was dissolved in alcohol and some strong hydrochloric acid added, by which a crystalline precipitate was formed, crystallising from hot acetic acid in beautiful red, glistening, flat needles. He was kind enough to give us a sample, and on examining it we found that water acted upon it in the same way as on our crystals.

In order to prepare a pure compound for analysis, a hot solution of aurin in acetic acid was saturated with hydrochloric acid gas; the colour of the liquid changed into a light yellowish red, and soon the compound separated out in glistening needles, which, even when perfectly dry, smell strongly of acetic acid. When exposed to the air they soon assume a steel-blue lustre, and gradually crumble into a reddish brown crystalline powder. The same properties are shown by the crystals obtained from acetyl chloride and those obtained from Mr. Lowe. When heated to 110° in a current of dry air they gradually lose all the acetic acid, which plays the part of water of crystallisation, and assume a dull red colour.

On passing hydrochloric acid gas into an alcoholic solution of aurin, similar but smaller needles are formed, containing alcohol, which is given off at 100° . The dull red residue can, like the preceding one, be heated to 100° in a current of dry air without losing hydrochloric acid, which only begins to escape at 200° .

Analysis of these compounds showed that the dried substance consists of $C_{19}H_{14}O_3 \cdot HCl$, while the crystals obtained from an acetic acid solution have the composition $C_{19}H_{14}O_3 \cdot HCl + 2C_2H_4O_2$, and those from alcohol $2C_{19}H_{14}O_3 \cdot HCl + 3C_2H_6O$.

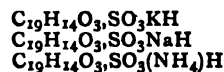
When sulphuric acid is added to a hot alcoholic solution of aurin, small red needles are formed on cooling, which consist of $(C_{19}H_{14}O_3)_2SO_3H_2 + \text{alcohol}$. Under the same conditions an acetic acid solution yields fine prismatic crystals or flat, very glistening needles, which are an acid sulphate, its formula being $C_{19}H_{14}O_3 \cdot SO_4H_2 + \text{acetic acid}$.

We have also prepared a nitrate which is readily formed and crystallises well, but have not analysed it yet.

In our first communication to the Chemical Society we described a compound of aurin and sulphur dioxide, which is easily obtained in bright red crystals by passing sulphur dioxide into a saturated alcoholic solution of aurin. Our former observation, that this body contains water but no alcohol, we found confirmed; on heating it decomposition

easily takes place, pure aurin being left behind, but it appears to be quite stable when exposed to the air, and even on heating it with water no sulphur dioxide is given off, but a drop of sulphuric acid added to the mixture is sufficient to evolve the gas abundantly. Aurin sulphite has the composition $(C_{19}H_{14}O_3)_2SO_3H_2 + 4H_2O$.

As we have already shown, aurin forms very characteristic compounds with the acid sulphites of the alkali metals, which, in accordance with the newly established formula of aurin, must now be written as follows:—



We have also found that rosolic acid, or the next higher homologue of aurin, forms compounds with mineral acids which crystallise well. Being, therefore, a base like aurin, we think its name ought to be altered, and, as it has only been obtained from rosanilin, propose for it the name *rosaurin*.

ON THE INFLUENCE OF CHLOROFORM ON NITRIFICATION.

By OTTO HEHNER, F.C.S., F.L.C.

NITRIFICATION, according to Schlösing, Muntz, and Warington, is prevented by chloroform.

The following experiments tend to prove that this is the case only when a relatively large amount of chloroform is employed, small quantities not only *not* preventing nitrification, but actually favouring the development of bacteria and the reduction of nitric acid into ammonia.

In a paper read before the Society of Public Analysts I have shown that on keeping ordinary drinking-water for some length of time the nitrogenous matter breaks up with the formation of ammonia, which in its turn gradually changes into nitric acid.

Attempting to arrest these changes by the addition of chloroform to polluted water, I was struck with the rapid development of bacteria in those samples which had received but small additions of chloroform, the samples becoming quite turbid and opalescent.

To $2\frac{1}{2}$ litres of a water polluted with putrid urine, yielding per 100,000 parts—

Free ammonia	0.0358
Albuminoid ammonia	0.0205

0.5 c.c. (2) and 1 c.c. (3) were added. After the lapse of three weeks sample 1 (without chloroform) showed—

Free ammonia	0.0007
Albuminoid ammonia	0.0065

Whilst (2) yielded—

Free ammonia	0.0992
Albuminoid ammonia	0.0480

And (3)—

Free ammonia	0.0109
Albuminoid ammonia	0.0659

Both (2) and (3) were very turbid when last analysed.

To samples of 8 litres each of another water, containing some putrid urine, 2 and 5 c.c. of chloroform were added. On January 18, 1878, the day on which the experiments were commenced, the samples showed—

Free ammonia	0.0239
Albuminoid ammonia	0.0430

On February 11 the sample without chloroform had changed to—

Free ammonia	0.0436
Albuminoid ammonia	0.0244

Whilst the water with 2 c.c. of chloroform contained—

Free ammonia	0.2017
Albuminoid ammonia	0.0208

And that with 5 c.c.—

Free ammonia 0.0227

Albuminoid ammonia 0.0404

The bottles containing the samples were kept freely exposed to daylight.

5 c.c. of chloroform had therefore prevented all change, whilst 2 c.c. had been quite ineffectual, the free ammonia having risen tenfold, doubtless at the expense of the nitrate contained in the water.

During the first week the albuminoid ammonia had somewhat increased in the latter sample.

Chloroform in small quantities is therefore rather a food than a poison to bacteria.

COMPOSITION OF THE GAS WHICH ISSUED FROM ONE OF THE SHAFTS OF THE ABERCARN COLLIERY.

By J. W. THOMAS.

It will be remembered by those who have read the reports of the terrible explosion which occurred at Abercarn on September 11 last, that in consequence of the coal having taken fire close to the main shaft at Abercarn and in the workings, and a direct communication between the upcast and downcast being determined by the explosive force which destroyed the parting doors, it was soon discovered that it was advisable to flood the colliery in order to quench the fire. It will be remembered, too, that the colliers who were fortunate enough to escape with their lives found their way out by the main winding shaft. The Abercarn Colliery has three shafts or pits—two downcasts and one upcast. The main downcast and upcast, which are 22 yards apart, are situated at Abercarn, and the other downcast is at Cwm Carn, a distance of a mile and a quarter in a direct line, and two miles underground from the upcast. As soon as the Abercarn portion of the workings was filled with water, the gas issuing from the coal and some of the products of combustion from the burning of the coal, and possibly more or less of the products of complete and incomplete combustion incidental to the explosion, were driven in the direction of the Cwm Carn pit, toward which the measures rise about 400 feet or a little over 2 inches per yard. After the flooding of the colliery was completed on September 16, when over 40 feet of water were in the two shafts at Abercarn—a large volume of gas, varying from 500 to 1500 cubic feet per minute, was evolved until the water was cleared out (about the middle of November) sufficiently low to admit of a current of air being drawn in the direction of the upcast shaft. On September 20th last I collected some of the gas issuing from the Cwm Carn shaft, which was covered with turf with the exception of a hole in the centre occupied by an iron pipe 11½ inches in diameter, through which the gas escaped. On both occasions the gas was collected in glass tubes previously fashioned for sealing off the ends before a blowpipe flame—a double acting syringe being employed to cause the gas to fill the tubes by displacement. The analysis of the sample of gas collected September 20 is marked No. 1. At the time of collecting it 500 cubic feet per minute were evolved, the temperature being 58° F. On October 9, I again collected gas, the composition of which is given in No. 2—about 1200 cubic feet of gas per minute were escaping at the time.

Composition of the Gas in 100 Parts.

	No. I.	No. II.
Carbonic anhydride ..	2.54	2.43
Air { Oxygen	2.73	none
Nitrogen	10.32	—
Marsh-gas	49.11	74.63
Hydride of ethyl	0.95	0.81
Nitrogen	34.35	22.13
	100.00	100.00

The composition of the hydrocarbons present, other than marsh-gas, resembled hydride of ethyl, but the analytical volumes did not agree well in No. I. On comparing No. I. gas with No. II. it is evident that the former contains a considerable proportion of nitrogen derived from atmospheric air, the oxygen of which had been used up to support the combustion of the fire-damp or the coal which was on fire. Owing to the plentiful supply of air which reached the burning coal previous to the colliery being flooded, it is possible that little if any carbonic oxide was generated, and the absence of this gas in No. I., in which it was carefully sought, seems to show that the incomplete products of the combustion of the explosive mixture (the fire-damp which gave rise to the explosion) had either in part been removed or so diluted with other gases that the carbonic oxide was present in quantity too minute to be determined in the process of analysis. The composition of No. II. is interesting. This consists doubtless of a mixture of blower-gases which found their way into the galleries through the channels and cracks in the strata, and the gases evolved from the working face or other exposed coal. The latter gases would be naturally of similar composition to those which escape under ordinary conditions of pressure and temperature from a lump of coal exposed to the atmosphere. In previous papers "On the Gases Enclosed in Coal," *Chem. Soc. Journ.*, 1875-6-7, I have pointed out on more than one occasion the curious fact that nitrogen escapes from coal in a vacuum at a proportionally quicker rate than any other gas. The flooding of the mine entirely cut off the supply of air and consequently no nitrogen could find its way into the galleries, admitting the possibility of the oxygen being used up. It is evident, therefore, that the percentage of nitrogen in No. II. is high, and that this circumstance points to the probability of its escaping from coal under ordinary conditions in the same manner as it does in a vacuum. Before No. II. sample was collected it will be seen from the analysis that all the air had been swept out of the workings by diffusion with the gases evolved from the coal, and long before the expiration of the nineteen days which elapsed between the collection of No. I. and No. II. all the products of combustion would have been removed. During the nineteen days an average of more than 1,000,000 cubic feet of gas was evolved each day, equal to the cubic contents of a large gallery or heading ½ miles in length. My thanks are due to Mr. Pond, of Abercarn, for the facilities afforded me in the collection of the gases and for his kind assistance.

The Laboratory, Cardiff, January 6, 1879.

ANALYSIS OF POPPY PETAL ASH.

By C. J. H. WARDEN, F.C.S.

ANALYSES of the organic constituents of poppy petals have been made by several chemists, but poppy petal ash has not hitherto, as far as I am aware, been examined. Before giving results of the analysis of the ash, a few remarks on the use to which poppy petals are put in the Government Opium Factories in Bengal may be interesting.

In the manufacture of opium for the China market—"Provision Opium"—it is customary to envelope the soft opium in a shell composed of poppy petals technically termed "leaves."

The petals are usually collected by noon on the third day of the flowers' expansion, and Mr. J. Scott thus describes the process:—"They clasp the petals at the base with the forefinger and thumb, gently drawing them upwards, and tightening them over the apex of the capsules, when the matured petals at once disarticulate or detach themselves. There is thus no tearing or straining of the petals, and no loss of juice from the disarticulated surfaces."

* "Manual of Opium Husbandry," by J. Scott.

The petals are then made into round "leaves" by spreading them on gently heated earthen or iron saucers, covering them with a moist cloth, and applying firm pressure by a damp pad. "Leaves" vary in diameter from 6 to 12 ins., and in thickness from 0.5 to 0.025 of an inch.

At the Patna Opium Factory 10 ozs. 94 grs. avd. of "leaves" are used for making the shell of each cake. The "leaves" are agglutinated round the opium by a 50 per cent mixture of opium in water.

Mr. Scott calculates that the annual consumption of poppy petals is upwards of 16,000 maunds (one maund equals 100 lbs. troy), to compose which the entire petals of no less than 4,710,400,000 flowers are required! During the season 1869-70 the Bengal Government spent £10,235 on leaves for the use of one factory. From the above short account of "leaves" it is evident that the poppy petal is an article of considerable commercial value.

The ash was prepared for analysis in the usual way, and was of a light grey colour, and effervesced with acids. An aqueous solution gave indications of the presence of SO_3 , P_2O_5 , Cl, and K. All the determinations were by gravimetric methods:—

100 Parts of the Ash Contained.]

Ferric oxide, Fe_2O_3	3.0536
Aluminic oxide, Al_2O_3	0.9701
Magnesian oxide, MgO	4.4311
Calcic oxide, CaO	8.4711
Potassic oxide, K_2O	32.9926
Potassic chloride, KCl	9.7078
Sodic chloride, NaCl	0.9537
Sulphuric anhydride, SO_3	3.0429
Phosphoric anhydride, P_2O_5	4.4331
Carbonic anhydride, CO_2	5.4662
Silicic anhydride, SiO_2	109.551
Sand	14.5881
Charcoal	1.1662

100.2316

Percentage of the ash after deduction of the unessential constituents, carbonic anhydride, sand, and charcoal:—

Ferric oxide	3.8647
Aluminic oxide	1.2278
Magnesian oxide	5.6082
Calcic oxide	10.7214
Potassic oxide	41.7569
Potassic chloride	12.2867
Sodic chloride	1.2071
Sulphuric anhydride	3.8513
Phosphoric anhydride	5.6107
Silicic anhydride	13.8652

100.0000

na Opium Factory, September 9, 1878.

IS CONTAMINATED WATER INJURIOUS FOR DOMESTIC PURPOSES?

By SERGIUS KERN, M.E., St. Petersburg.

THE St. Petersburg town council organised a committee for the discussion of the following question:—About seven miles from the town a new cemetery was opened some years ago. A small river runs from this place into the Neva, some eight miles above the town. It was found that this small river is entirely contaminated by the decomposition of the bodies. Owing to this inconvenience many propose to close the cemetery. The discussions which took place in the sitting of the committee on November 23, 1878, are rather interesting, as the sanitary improvements of the town are still considered secondary questions.

M. Borodin, Professor of Chemistry, thinks that water

containing a certain amount of putrescent substances cannot have a great influence on the health of the inhabitants of the town, and remarked that even in science this question is nearly an open one, as it has been only partly discussed. Indeed, it is a question if contaminated water, used for domestic purposes, can do any harm. The Professor remarked, also, that sometimes, for instance, in consuming various cheeses we introduce at one time a quantity of putrescent matter into the stomach, which would not be detected in the annual amount of water consumed, containing a large quantity of organic matters. The analyses of the Neva water (by Professor Trapp) show that it cannot be called bad water:—

In 1 Cube Metre.

Inorganic matter	32.25 grms.
Organic matter	23.00 "
CaCO_3	15.25 "
MgCO_3	7.40 "
NaCl	1.60 "
Na_2SO_4	2.40 "
K_2SO_4	1.30 "
FeCO_3	1.20 "
Al_2Cl_6	2.60 "
SiO_2	0.50 "

M. Bertenson could not agree with the Professor. He remarked that oxygen is very important in water, and certainly the more organic matter is present in the water the less it contains of free oxygen.

The committee decided to inspect the cemetery, and take some water for analysis from the small river, and then agreed to inspect also one of the bodies which was buried the first on this ground. The conclusions of the committee will be of some interest, as there is not much agreement between the members concerning the question discussed.

THE URIC ACID GROUP.

By SAMUEL E. PHILLIPS.

As Mr. P. T. Main has advanced some new views of urea, uric acid, and their derivatives, it may interest some students of chemical philosophy if I give the substance of a paper read before the British Association of 1877, "On the Principle of Uric Acid Genesis."

"As the vast and complex researches of Liebig, Strecker, Baeyer, and others have now culminated in the studies of M. Grimaux, it is thought to be high time to trace herein the plain indications of law, order, and symmetry in lieu of empirical confusion." The author refers to a paper on the constitution of uric acid (CHEM. NEWS, vol. xxxii., p. 209), and now maintains that a clear view of the chemical laws involved in -2HO , under varied conditions, will supply the key to unravel the endless empirical confusion.

Starting with eight acids, it is maintained that under the combined influence of heat, time, and pressure that all these, and some others, act similarly upon urea, and condense with successive eliminations of -2HO , in proportion to the strength or duration of the applied conditions:—

Hydromellitic acid (?)	$\text{C}_4\text{H}_2\text{O}_2\text{O} \cdot \text{HO}$
Glycollic acid	$\text{C}_4\text{H}_3\text{O}_4\text{O} \cdot \text{HO}$
Glyoxalic acid	$\text{C}_4\text{H}_1\text{O}_4\text{O} \cdot \text{HO}$
Glyoxylic acid	$\text{C}_4\text{H}_3\text{O}_6\text{O} \cdot \text{HO}$
Unknown acid	$\text{C}_6\text{H}_7\text{O}_2\text{O} \cdot \text{HO}$
Pyruvic acid	$\text{C}_6\text{H}_3\text{O}_4\text{O} \cdot \text{HO}$
Malonic acid	$\text{C}_6\text{H}_1\text{O}_4\text{O} \cdot 3\text{HO}$
Tartronic acid	$\text{C}_6\text{H}_7\text{O}_6\text{O} \cdot 3\text{HO}$

That with these, eight ureide series may be projected, embracing very much that is already known, and presenting

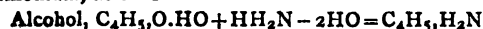
some blanks, which may easily be attained in the future, and that these eight series may be thus epitomised, assuming 1 equivalent of acid in each case:—

- + Urea $-2\text{HO} = (\text{CO})_2\text{RH}_3\text{N}_2 = \text{the R urea (a)}$
 $-4\text{HO} = \text{CyRH}_3\text{N} = \text{the cyanamid (b)}$
 +2 Urea $-2\text{HO} = (\text{CO})_4\text{RH}_7\text{N}_4 = \text{the di-urea (c)}$
 $-4\text{HO} = (\text{CO})_2\text{CyRH}_5\text{N}_3 = \text{the pseudo-uric acid (d)}$
 $-6\text{HO} = \text{Cy}_2\text{RH}_3\text{N}_2 = \text{the ultimate (e)}$

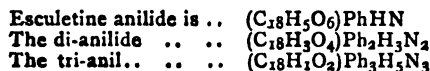
It is remarked that these uric acid *ultimates* are not strictly such, and the two extensions of variety are excluded from the brief terms of the paper:—First, certain obscure cases where these radicals are affected by reaction with ammonia -2HO , as in hydrazulmoxin and hydrazulmin; secondly, the more complex forms of ureide condensation obtained by M. Grimaux, which all, nevertheless, prove to be grand illustrations of the same definite laws.

It follows that all the (a's) are normal ureas, though often associated with confused names in their empirical discovery; that all the (b's) are R cyanamids, though not at all recognised as such, and some of them are called ureas; that the ureides or di-ureas have three stages ending in uric acid and its other analogues; that the law of elimination has three aspects:—

1. When hydrates condense into each other, or into an ammonium, as thus—

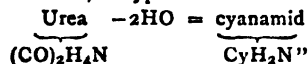


2. When successive elements of ammonia are super-added to the condensation, each step is attended with an additional -2HO by derivation from the radical when no hydrate subsists. A wonderful regularity may be traced in the glucuride researches of M. Schiff, by which for illustration—



And a recent research of M. Ladenburg's, where hydrates are "condensed into toluylene diamine, affords a striking illustration, the more so, as given like the former, on empirical and independent grounds.

"3. When ureas condense into ureides, the tendency, after eliminating -2HO for each equivalent of acid or hydrate, is for each $(\text{CO})_2$ to change into cyanogen with elimination of 2HO , as typified in the well known case—

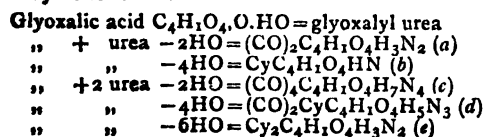


I am much pleased to find that I am not alone in entertaining the cyanamid character of uric acid; but the idea of including urea in the same category, as a derivative of uric acid, is certainly a confused and illegitimate sketch of hypothesis.

The ammoniacal character of urea is established beyond question; its amide dedoublements and their basic characters admit of no other explanation.

M. Dreschel, in obtaining several varied cyanamids, notates one as "the hydrochlorides." It is therefore clear that he might have obtained the hydrate. Besides, hydrates of dicyanamid and others are known, but surely no one expects to find them ureal.

Mr. Main falls into a mistake, which his notations ought to have corrected; but, alas! they are only fitted to mislead. Allantoin is not a derivative of "glyoxylic acid," as may be seen thus:—



(b) As urea -2HO becomes cyanamid, so glyoxalyl urea -2HO becomes glyoxalyl cyanamid.

(c) Is the hypothetical first stage of the di reaction.

(d) The second stage is *allantoin*, and the formula of production is precisely that given by the discoverer, M. Grimaux. It also corresponds with M. Grimaux's *Pyruvil* in the pyruvic series, or with my pseudo-uric acid in the tartronic series.

(e) Is the ultimate mis-called "mycomelic acid" and the perfect analogue of uric acid.

One word more as to the *bizarre* character of these misleading notations.

The discoverer of "melid acetic acid" thought it was an aceto body, and that it was an acid, and hence the terminal (COOH) so fashionably characteristic of acid types! Of course the pretence is that these structural or graphic portraits are intended to delineate certain features of genetic or analytic behaviour in the compounds. The plain fact is, however, that it is a mischievous device of wanton hypothesis, and hence our exposure of the true nature of that body, as containing no acetic acid in its genesis, and no acid behaviour in its derivatives. It is a substituted *mellamine* base! (See CHEMICAL NEWS, vol. xxxiv., p. 13.)

When M. Grimaux speaks of glyoxylic acid he means glyoxalic acid, and his notations correspond with his remarkable synthesis of allantoin: but Mr. Main, rightly estimating the total elements of glyoxylic acid, gives allanturic acid as one of its derivatives, which is a double mistake. First, as a matter of chemical fact; and, secondly, as a matter of notational display.

Allanturic acid is "glyoxalyl urea." Why, then, justify its early misnomer by an *acid* notation ending in (COOH)

Glyoxylic acid—



Allanturic acid—



And why notate it with cyanogen as a constituent?

Under the derivatives of tartronic acid we have—

Dialuric acid—



Violuric acid—



(a) is tartronic urea, $(\text{CO})_2\text{C}_6\text{H}_7\text{O}_6\text{H}_3\text{N}_2$. Both (a) and (b) are given with the acid terminations to correspond with tartronic acid; but it is quite overlooked that (b) belongs to another acid series, for it is known to be a "malonyl urea."

How, then, does it happen that so many amides should be called acids? The answer is plain, and the parentage is clearly traceable to a mistaken hypothesis of the early discoverers, which is grossly perpetuated to this day. They noticed among these bodies a tendency to combine with or to assimilate metal ingredients, and the notion was that the relation subsisting was that of acid and base.

It is, of course, familiar that many hydrocarbons have a tendency which is notably typified in acetylen; and we now find that the same tendency is frequently evinced among amides; but the wonder is that modern chemists should be so slow in establishing a just discrimination.

Were we to imitate Mr. Main's ureal hypothesis, and contend that ammonia was an acid, and that the amides of Zn, Cu, or K were its basic salts, the idea would necessarily be scouted; but the same principle is largely and widely indulged in throughout the modern types of organic chemistry; and we constantly read of amido acids, and see their terminal (COOH) graphically displayed, while the context in so many cases reveals the fact of their *basic* characters, as seen in their "hydrochlorides," their sulphates, nitrates, and other salts.

I care not to follow Mr. Main when he seeks to paint the lily, and to improve his *bizarre* notations by atomicity equations and "more symmetrical" portraits of the atomic dispositions. But the body selected for all this display is wisely chosen from the least understood of chemical compounds, viz., the amorphous substance called "hydrazulmin."

It may or may not be a definite chemical body; but as I have done my best, in a known and legitimate way, to arrive at some faint *a priori* understanding of its genesis and relations in a yet unpublished "Study of Ureide Derivatives," it may be well to compare notes as to our methods of investigation. One is playing with shillies, and can place them in any position, only controlled by certain laws of the game, such as the atomicity rules, &c. The other is rather a student of the actual chemical forces involved, and has comparatively a very limited number of pieces. He can only place on the board definite or well known radicals, and these only in certain limited modes of relationship, while from an infinitude of "residues" he is utterly debarred.

With one it is an easy matter to build up on paper a melid-acetic acid, or an allanturic acid, with wrong materials; with the other these feats are simply impossible. One is so confidently assured that he can open out the condensed types to graphically exhibit the internal play of atomicities, and go yet further in the Frankland school to "more symmetrical" pictures. The other has to retire in all humility, simply confessing inability to either execute or understand such works of art.

"We have referred to the uric acid analogues as *ultimates*. That term, however, admits of some qualification, because by another mode of reaction it is possible to reduce the radicals of the varied acids, and in that respect to realise more ultimate compounds.

- "Mycomic acid," $Cy_2C_4H_2O_4H_3N_2(a)$
 $(a) + H_3N - 2HO$, hydrazulmoxin,
 $Cy_2C_4H_2O_4H_2(NH_2)_2N_2(b)$
 $(b) + H_3N - 2HO$, hydrazulmin,
 $Cy_2C_4H_2OH(NH_2)_2N_2(c)$

It only remains to notice that this hydrazulmoxin is identical or isomeric with Mr. Main's "azulmic acid."

That in each of these stages of reaction one equivalent of ammonia is added is a chemical fact beyond question; and that each stage implies a theoretical tendency to $-2HO$ is a principle I have tried hard to exemplify, and need not here repeat the grounds, while gladly referring to the actual facts of analysis in confirmation.

PROCEEDINGS OF SOCIETIES.

NEWCASTLE CHEMICAL SOCIETY.

General Meeting, November 28, 1872.

Mr. R. C. CLAPHAM, President, in the Chair.

THE minutes of the previous meeting were read and confirmed.

The following were elected members:—Mr. Walter Weldon, Dr. Bernard Mohr, Mr. John Watson, Mr. Thos. Campbell, Mr. John Hope, jun., Prof. R. W. Atkinson, Mr. Mayfield, Mr. J. W. White, Dr. H. E. Armstrong, Mr. T. Crawford, Mr. W. G. Strype.

The following name was read for the first time:—Mr. John Cliff, Runcorn.

THE PRESIDENT—Our first business, gentlemen, is to proceed to the discussion on Mr. Pattinson's paper, "On the Quality of the Small Coal used on the Tyne." It is a very valuable paper, especially as regards the amount of sulphur contained in the ash. I do not remember having before seen results set forth to show both the vola-

tile sulphur and the sulphur contained in the ash. Perhaps it might be interesting to the members if Mr. Pattinson would explain how the amounts of these latter are determined.

MR. PATTINSON—It is very quickly done. The total sulphur is determined in the usual way, by oxidation; then the amount remaining in the ash of the coal is determined, and the difference of these two amounts gives the volatile sulphur. The sulphur in the ash is retained by virtue of the lime which the ash contains, which probably existed originally as carbonate. I may perhaps mention here the method used to determine the total sulphur in the coals. The old method, by heating with strong nitric acid and evaporating to dryness, is very incorrect, never yielding more than a fraction of the sulphur really contained in the coal. The method by deflagration with nitre and salt is very correct when carefully done, but it requires great care, both in the deflagration and in the subsequent precipitation of the sulphur as barium sulphate. The method now used, however, is very much simpler than any of the old ones: a weighed quantity of the coal or coke (about 20 grains) is mixed with one and a half times its weight of slacked lime, moistened slightly with water, and then heated in a muffle at a red-heat. The carbon burns off, and the sulphur is converted into calcic sulphate, which is then dissolved in hydrochloric acid, and the sulphuric acid precipitated in the usual way.

MR. H. L. PATTINSON—What is the reason that the method of treating with nitric acid is unsatisfactory?

MR. J. PATTINSON—Probably a portion of the pyrites is covered and protected by the coal, even when it is very finely pulverised.

THE PRESIDENT—If no member has any more remarks to make we will proceed to the discussion on Mr. Dunn's paper, "On Indicators in Alkalimetry." I do not know if Mr. Dunn has anything further to add on the subject of his paper.

MR. DUNN—I have not done anything more in the matter since the paper was read. I may just mention, however, that aurin does not keep quite so long as litmus. It will, however, keep for two or three months without alteration, and the preparation of the solution is so easy that it can be made afresh every two or three months with very little trouble.

THE PRESIDENT, in calling on Mr. Cookson to read his paper, "On Rozan's Process for Desilvering Lead," said, I had the pleasure of going down to Messrs. Cookson's works, at Willington, a few weeks ago, and I think I have not for a long time been in any works which were more interesting, both from a chemical and a mechanical point of view. The handling of large quantities of lead is done almost entirely by hydraulic machinery, which I have no doubt saves a large amount of money formerly spent in wages for hand labour. The paper to come before us to-night is one of very great interest, and perhaps, if we say anything about it now, it must be to express a kind of regret that a patent is likely to come into operation which may do away with the classical process of Mr. H. L. Pattinson—a process for which the North of England has long been justly celebrated.

"On Rozan's Process for Desilvering Lead," by Mr. COOKSON.—In attempting to describe the leading features of the Rozan system of desilvering, I must ask you to bear in mind that in principle it is in most respects identical with the well-known process of the late Mr. H. L. Pattinson, always excepting that in the former case cranes and steam supersede the more expensive system of manual labour employed by Mr. Pattinson. In both cases, however, the whole working depends on the fact that as melted lead cools the first-forming crystals contain less silver than the portion remaining liquid, and that in the usual way of working by thirds this exists to the extent that when two-thirds of a given quantity is in crystals and one-third liquid, the two-thirds will actually contain only half the quantity of silver that the one-third or lesser quantity does. In the Rozan system steam is

used with a double purpose, one of which is mechanical and the other chemical. The mechanical effect is to boil up and violently stir the melted lead, thus preventing any setting on the surface; and this boiling of the steam through the lead as it cools causes a regular crystallisation, and an easy and perfect separation of the liquid from the crystals when the proper stage is reached. The chemical effect is due to the oxidising action of the steam on the antimony, copper, iron, arsenic, and other extraneous metals, which, being converted into oxides, are either skimmed off or carried away with the escaping steam, and afterwards allowed to settle in the condensers. This chemical action fulfils so good a purpose that the Spanish rich silver leads are desilverised without any previous calcination, and an unusually pure market lead is produced from them. Even in the case of Greek lead only a partial calcination is required, as experience shows that a portion of antimony, equal to about $\frac{1}{4}$ per cent, existing in the original silver lead, is an advantage, inasmuch as this small quantity of antimony has a remarkable effect in reducing the quantity of lead oxides formed. Without absolute proof it may be premature to speak of any chemical action, as it is possible the whole of the oxidation is due to the mechanical exposure of the molten impure lead to the influence of the atmosphere, assisted by the air carried into the pot by the steam; but I am strongly of opinion that a chemical action actually takes place. The leads most suitable for the steam process are of a similar class to the already-mentioned Spanish rich silver leads, containing from $\frac{1}{4}$ to $\frac{1}{2}$ per cent of foreign metals, of which antimony generally constitutes fully two-thirds of the quantity. In the old Pattinson process those extraneous metals are eliminated as far as possible by a previous calcination, the charges sometimes requiring forty-eight hours in the calcining furnace in order to fit them for the pots. In the Rozan system, however, this calcination is dispensed with, and the lead as it comes out of the smelting furnaces is used direct by the desilverising apparatuses.

Having described the working of the plant, I will put before you as shortly as I can the advantages and defects of the Rozan as compared to the Pattinson system as carried out in our works for a great number of years. The advantages the Rozan system possesses are—

1. The entire saving of the cost of calcining all ordinarily hard leads, and in the case of extra hard leads, such as Greek lead, a very large saving.
2. A cost for labour not exceeding one-fifth.
3. A cost for fuel of about two-fifths.
4. A saving of one-third in the oxides produced, which advantage any lead manufacturer will fully appreciate.

Its defects are—

1. A large capital outlay.
2. A constant expense in repairs and renewals.

The repairs are only a small matter, but until lately the cost of renewal of pots was a very serious and heavy item: it is one which by observation and care we have been able very considerably to reduce. In 1876 pots cost us over 3s. per ton of market lead; in 1877 it came to 2s. 2d.; and this year, so far, the cost has been further reduced to 1s. 4d. per ton.

On the whole, the advantages are very considerably greater than the defects of the Rozan system; and I can confidently assert that our firm has every reason to be satisfied with having adopted it.

The following is a table showing silver assays of twelve crystallisings taken from an average of 350 operations:—

Ounces per ton of lead.

570	315	202	112½	62	33½	19½	10	5	2½	1½	14 dwts.
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The PRESIDENT—Would any member like to ask Mr. Cookson any questions about the paper? Mr. Morrison has referred to condensation, and it struck me that that was a point which, at the present meeting, Mr. Cookson

might give a little more account of. When I saw Mr. Cookson's works the condensation appeared very perfect and free from escape.

Mr. COOKSON—It is not a point of much importance whether the condensers are air-tight or not, because we have a draught at the far end, so that there is always an inward suction. In practice we find that in the last chamber there is little or no settling. I have sometimes opened the door and gone in, but could never observe anything except steam in the state of fog or cloud. The oxides of antimony and copper are partly skimmed off the melted lead as dross, and partly condensed as fume in the chambers. The fume near the furnaces contains a larger quantity of these oxides, and as it gets further from the furnace it approaches more and more nearly in composition to pure oxide of lead, thus confirming an old dictum of lead-smelters, that oxide of lead is more difficult to condense than the oxides of the accompanying metals. One thing I did not mention, which has indeed nothing to do with the subject of the paper, but is simply a mechanical arrangement—our system of cranes is very perfect. From the time when the crude lead enters our works to the time when it makes its exit as "market lead" it is hardly ever touched by hand, but lifted from place to place entirely by hydraulic cranes, which costs very much less than if it were lifted by hand.

Mr. GLOVER—In blowing the steam through is there any chemical decomposition of the steam, or is the action simply mechanical? Are the copper and antimony oxidised by the steam, and the oxides volatilised, or carried off mechanically?

Mr. PATTINSON—To a chemist the most interesting point in the process is that which has just been mentioned by Mr. Glover. I am inclined to think that at such a temperature the copper would not be volatilised. Could Mr. Cookson tell us the composition of the fume condensed near the crystallising pot? I think that as the oxides of copper and antimony accumulate on the top, they are most likely to be mechanically carried away to the condensers. The question is, is the steam decomposed, is hydrogen produced, or is the oxidation due to the atmosphere?

Mr. COOKSON—I think it is due principally to the free oxygen which is always contained in small quantity in the steam, and that the oxides are then carried away by a purely mechanical action.

Mr. B. S. PROCTOR—Have you ever had any trace of hydrogen?

Mr. COOKSON—I have put my nose into the condensers, thinking that if hydrogen were produced there would be a certain quantity of arseniuretted hydrogen formed, but I have never smelt any. There is a trace of sulphur found in the condensers, but I never smelt any sulphuretted hydrogen.

Mr. H. L. PATTINSON—Have you ever tried air instead of steam?

Mr. COOKSON—It is included in the patent, but it increases the quantity of oxide of lead formed very greatly.

Mr. GIBB said, it appeared to him that the action of the steam was purely mechanical, not chemical, and that the process was merely the Pattinson process with improved mechanical arrangements.

The PRESIDENT wished to remind Mr. Gibb that most of his objections had been already answered by Mr. Cookson in the paper itself.

Mr. COOKSON—I so far agree with Mr. Gibb that I often call the process "Mechanical Pattinsonisation." The steam may perhaps be described as a vehicle for conveying the oxygen of the air in limited quantity to the lead.

Mr. PATTINSON—Or a valuable means of getting the mechanical effect of air with a smaller proportion of oxygen.

Mr. MORRISON—The steam protects the lead?

Mr. COOKSON—No; for if we work soft leads we find, with all our care, that the quantity of oxides produced is largely increased.

The PRESIDENT—I will remind the members at this stage that Mr. Cookson's paper will come on for discussion at another meeting; meantime I propose a very cordial vote of thanks to Mr. Cookson. The description which he has given is very interesting, and we are all much indebted to him for such an able paper on this important question.

The motion on being seconded was carried unanimously.

NOTICES OF BOOKS.

Fourth Annual Report of the Commissioner of the Imperial Mint, Osaka, Japan. Hiogo: Hiogo News Office.

This report contains nothing of special scientific interest. It appears that the foreign staff of the Japanese mint now consists of two persons only, the chemist, Mr. W. Gowland, F.C.S., and the engineer, M. R. MacLagan, M.I.M.E., all the other posts being now filled by natives. The sanitary condition of the Mint department does not seem satisfactory; 411 cases of illness and 11 deaths out of a total of 602 men in their best years are remarkable, especially as none of the deaths were due to accidents. How much of this is due to the climate and how much to the nature of the employment cannot be known, especially as we can find no mention of the number of men in each department.

The Magic Lantern Manual. By W. J. CHADBURN. With 100 Illustrations. London: Warne and Co., 1878.

POOR Artemus Ward used to call himself a "jokist," so Mr. Chadburn takes liberties with Her Majesty's English and calls himself a "Lanternist." Whether he will succeed in adding a new word to the dictionary or not we cannot venture to prophesy, but in any case he has presented his brother "lanternists" with a capital little book on the magic lantern. He first of all describes the ordinary old fashioned oil lantern and its objectives, and then proceeds to give us an account of the Sciopticon, an American invention, which ranks half way between the old fashioned instrument and the oxyhydrogen and electric lanterns. The pyrohydrogen and magnesium lanterns are also described, as well as all the accessory instruments connected with them. Clear directions are also given for the preparation of the slides, both by hand and by photography, whether by the silver or carbon process. The part, however, which will most interest our readers is that on scientific projections. For exhibiting diagrams, optical effects, the decomposition of water, crystallisation, &c., Mr. Chadburn tells us that the Sciopticon will serve every ordinary purpose for class demonstration. This is a fact that is evidently but little known to science teachers, who always look on lantern demonstration as involving endless trouble and an enormous outlay. We cordially recommend Mr. Chadburn's little book to all science teachers and lecturers.

The Unity of Matter: A Hypothesis. (Die Einheit des Stoffes: Eine Hypothese). Von A. WALDNER. Zürich: Zürcher and Furrer.

THE author of this little pamphlet tells us in his preface that the hypothesis in question is the result of an arithmetical examination of the relations existing between the atomic weights of the so-called chemical elements. He undertook this investigation in June last (1878), but did not make his results public, as no facts were known to prove that these elements were capable of further decomposition. The recent researches of Mr. Norman Lockyer and his experiments on the transformation of the metals have induced him to publish his hypothesis, which they seem to confirm. He considers it possible that the

formulæ given below (in thesis 10) may be capable of still further simplification.

We give an abstract of the author's theses:—

1. All have arisen out of the so-called cosmic ether, the one primitive matter which fills infinite space.
2. This primitive matter, whose motions make themselves known by the phenomena of light, heat, electricity, and magnetism is not infinitely divisible.
3. The portion of the original matter, which after continued division resists all further comminution, is the primary atom.
4. All primary atoms have identical (regular) form and equally great mass (weight), and are amongst themselves perfectly congruent.
5. The cosmic ether, the initial state of all nascent bodies, is formed of primary atoms lying beside of each other without definite law.
6. In these primary atoms there inheres a power or tendency to combine with each other.
7. By these combinations of the primary atoms there arise molecules of new bodies different from the primary matter.
8. The combinations of the primary atoms take place according to the following law:—

$$\begin{array}{lll} (1.) & 2^1 & 2^2 & 2^3 \\ (2.) & 3^1 & 3^2 & 3^3 \end{array}$$

i.e., primary atoms unite together either by twos and threes, or by the third and fifth powers of two and three.

9. These six primary compounds form the basis of all known bodies, the molecules of the so-called elements being formed by combinations of such primary compounds.

10. These combinations take place simply according to the law of multiple proportions. If in series (1)—

$$2^1 = 2 = a; 2^2 = 8A; 2^3 = 32 = a$$

and in series (2)—

$$3^1 = 3 = b; 3^2 = 27 = B; 3^3 = 243 = b$$

the molecules of bodies considered as elements are formed as follows:—

$$\begin{aligned} a_2 b_2 &= 2 \times 2 + 2 \times 3 = 10 = \text{H.} \\ a_4 b_2 &= 2 \times 32 + 2 \times 3 = 70 = \text{Li.} \\ a_2 b_{10} &= 2 \times 32 + 10 \times 3 = 94 = \text{Be.} \\ a_1 b_4 &= 1 \times 32 + 4 \times 27 = 140 = \text{N.} \quad \&c. \end{aligned}$$

11. If the mass (weight) of the primary atom = 1 the above numbers show not merely the number of original atoms in the compound molecules, but also their molecular weights.

12. The conditions under which the original atoms combine together, and under which, inversely, compounds are resolved into their original matter are hitherto unknown, and are probably of a nature scarcely to be realised by the temperatures and pressures at the command of our present science and technology."

On this proposed law criticism would be obviously premature. But we do not see on what principle the author selects the factors by which a, A, and a, and b, B, and b are respectively multiplied.

CORRESPONDENCE.

A NEW TEST FOR ARSENIC.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxviii., p. 301, it appears that Mr. Otis Johnson claims the discovery that when potassium hydrate is allowed to act upon aluminium in the presence of arsenic and antimony, that AsH_3 is evolved without the evolution of SbH_3 . The above reaction is not new, having been discovered by M. Filhol, professor at Toulouse, prior to July, 1876. Prof. Filhol proposed the following, viz., if hydrogen is allowed to act

on arsenic by means of zinc or aluminium and caustic potash, and not by Zn and dilute sulphuric acid, as has always been customary in the Marsh apparatus, AsH_3 is readily evolved, but if tried with antimony compounds not the least trace of SbH_3 is observable. It is thus possible in the case of a mixture of As and Sb to disengage all the As and leave the pure Sb behind. It is also worthy of notice that when hydrogen is evolved by action of Zn, or still better Al, on potassium hydrate, if phosphorus be present, the colour of the ignited gas will be a beautiful green. The least traces of phosphorus can in this way be detected.—I am, &c.,

WILLIAM JOHNSTONE.

Analytical Laboratory, Lowther Hill,
Forest Hill, December 28, 1878.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 26, December 23, 1878.

Explosion of Deflagrating Matter.—M. Dupuy de Lôme.—The author examines into the causes of an explosion by which a M. Zédé had been severely wounded. The latter was endeavouring to find a compound which without exploding should be entirely resolved at the lowest possible temperature into gases and vapours, and which should serve as a motive power. For this purpose he employed a mixture of gun-cotton and of nitrate of ammonia. After finding the most suitable proportions he was studying in how far the speed of combustion, very slow in the open air, might be modified under increased pressure. On one occasion, when setting fire to the mixture contained in his apparatus, there occurred a violent explosion, attended by a flash of light. The tube, which had been tested up to fifty atmospheres, was shattered to pieces, and the experimentalist was seriously wounded. It would appear that a slight decrease in the orifice through which the gases escaped had changed the nature of the process from deflagration to detonation.

Magnetic Rotation of the Plane of Polarisation of Light under the Influence of the Earth.—H. Becquerel.—The author points out that the experiment described in M. Joubert's paper (*Comptes Rendus*, lxxvii., p. 984), is a reproduction of one which he had described on p. 1075 of the preceding volume.

Novel Phenomenon of Static Electricity.—M. Duter. A continuation of the controversy between the author and M. Govi.

Preparation of the Cobalto-cyanide of Potassium and of some of its Derivatives.—A. Descamps.—On pouring a cold solution of cyanide of potassium into chloride of cobalt there is formed a reddish brown precipitate of cobaltous cyanide. Care must be taken not to pass the limit of decomposition. This precipitate is kept at 0°, carefully washed in water, and then dissolved in a slight excess of cyanide of potassium at the same temperature. The liquid if diluted with alcohol deposits crystals of a deep amethyst blue. These are washed in alcohol to remove the excess of cyanide of potassium, and are then preserved in alcohol at 95°. This salt is unstable and soon turns red. If dissolved in a little water it gives a deep red solution, which yields several reactions. With acetate of lead it yields an orange-yellow precipitate of cobalto-cyanide of lead. With chloride of cobalt, absolutely free from nickel, it gives a deep green precipitate, the cobalto-cyanide of cobalt and potassium.

Action of Trimethylamin on the Sulphide of Carbon.—A. Bleunard.—These two substances react upon each other with some violence, forming sulphocarbamate of trimethylamin.

Biedermann's Central-blatt.

Heft 12.

On the Requirement and the Transformation of Matter in *Saccharomyces Mycoderma*.—A. Schulz.—Not suitable for abstraction.

Salicylic Acid as a Disinfectant.—Prof. Feser.—The author, in a prolonged series of experiments, found salicylic acid of no value, either as a prophylactic or in the treatment of putrid infectious disease, carbuncle, &c.

Revolution in Tanning.—Prof. Knapp proposes the use of a basic ferric sulphate instead of oak-bark or other tanniferous material. He adds to a boiling solution of copperas the quantity of nitric acid requisite for the peroxidation of the iron, and after the reaction is over adds more copperas. The hides are suspended in the cold solution at a suitable degree of concentration, and are ready in from two to four days.

Chemiker Zeitung.

No. 50, 1878.

According to Dr. Deite most of the French manufacturers of iodine who buy their weed-ash from the varec-burners obtain only 3'15 kilos. of iodine from a ton.

The yearly production of potash in France is estimated at 14,000 tons, 10,000 of which are obtained from beet-ashes and 1000 tons from the suint of wool.

The importation of American leather into Germany has increased from 23,738 cwts. in 1868 to 155,773 cwts. in 1875. A strong demand for protective duties has arisen among the German tanners.

A successful competitor at the late Paris Exhibition offers his bronze-medal as a contribution to the lottery, as it was awarded him without any examination of the quality of his exhibits.

A long list is given of the substances used—apparently in France—for "denaturing" common salt to be used for agricultural or technical purposes. Some of these additions must greatly increase the price.

Melted gallium dissolves aluminium even below 15°, forming liquid or semi-liquid amalgams which oxidise very slightly on exposure to the air, but decomposed water powerfully, the gallium being liberated in the form of metallic globules.

Dr. Kayser, after examining pigments such as white-lead, zinc-white, chrome-yellow, &c., finds that in addition to the heavy-spar, the quantity of which is stated by the seller and is allowed for in the price, gypsum is likewise present.

Prof. Schwalbe contributes a judicious essay on chemical nomenclature. He considers it unwise to attempt the introduction of new names for long-known compounds, and thus to carry us further and further away from the desired goal,—a universally intelligible terminology.

Doubts are expressed whether the electric light can compete with coal-gas from a commercial point of view.

The dangerous character of lead-compounds is shown by the fact that in the years from 1838 to 1847 no fewer than 3142 patients suffering from lead-cholic were admitted into the hospitals of Paris, although there were at that time only two white- and red-lead works in the city. Of these cases 112 proved fatal.

Plicque estimates the total yearly production of artificial ultramarine at 10,000,000 kilos.

Incrustation of Lead Pipes with Sulphide of Lead.—A hot concentrated solution of sulphide of sodium is

allowed to flow through the pipes for ten to fifteen minutes. They then appear as if coated within with a grey glaze, and water afterwards passed through them remains free from lead.

According to *Dingler's Polyt. Journal* the skin of the sting-ray (*Raja clavata*) is now used in place of isinglass for the clarification of liquids, e.g., beer.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 13, 1879.

Conversion of the Nitriles into Imides (Fourth memoir).—A. Pinner and F. Klein.—The authors here examine the action of hydrochloric acid and alcohol upon hydrocyanic acid, which gives rise to ammonium chloride, ethyl chloride, formic ether, diethyl-glyoxylic amide, and diethyl-glyoxylic ether. When the same compounds react upon cyanogen the result is a predominating quantity of oximide ether, and secondarily chlor-ethyl, formic ether, and urethan. The authors have also studied the action of hydrochloric acid and alcohol upon propionitrile.

On Butyl-chloral-cyanhydrate.—A. Pinner and F. Klein.—The authors study the behaviour of this compound with ammonia, urea, acetyl-chloride, concentrated sulphuric acid, hydrochloric acid with alcohol, and other compounds. They attach particular interest to the action of ammonia upon trichloroxy-valerianic acid.

On Naphthyl-phospho- and Arsenio-Compounds.—W. Kelbe.—An account of naphthyl-phosphorous acid, diethyl-naphthyl-phosphin, triethyl-naphthyl-phosphonium iodide, dinaphthyl-phosphinic, and naphthyl-arsinic acid.

Determinations of Specific Weights.—F. W. Clarke.—Not susceptible of useful abstraction.

Etherification of Primary Alcohols.—N. Menshutkin.—The author points out and rectifies certain errors in his paper, *Berichte*, x., p. 1728.

Action of Nitrous Acid upon Non-saturated Hydrocarbons.—Paul Tönnies.—In this preliminary communication the author mentions that if a saturated solution of potassium nitrite is brought in contact with a solution of the above compounds in glacial acetic acid, substances are formed which appear as addition-products of N_2O_3 .

Preparation of a Series of Magnetic Compounds of the General Formula RO, Fe_2O_3 or $R''Fe_2O_4$.—Karl List.—The author has obtained and examined the ferrates of lime, magnesia, manganese, zinc, nickel, copper, and lead.

On Cinchonin and Cinchonidin.—Z. K. Skrap.—To the former of these the author ascribes the formula $C_{19}H_{22}N_2O$, and considers that cinchonidin has exactly the same composition.

Remarks on the Previous Paper.—O. Hesse.—The author finds the amount of carbon in cinchonin decidedly higher than does Skrap.

Analysis of the Mineral Spring "Tenniger Bad" in Somvixer Tobel.—R. Meyer.—The water contains an unusual quantity of strontium.

Angelic Acids of Different Origins.—W. v. Miller.—Frankland and Duppa's methyl-crotonic acid is distinct from Neubauer's angelic acid. The author's is a new isomer, of the formula $C_5H_8O_2$.

Preservation of Drinking Water.—Hugo Schiff.—The author recommends salicylic acid for the preservation of drinking water and brine with 1 part per 1000 of purified carbon disulphide for the preservation of zoological specimens.

Oxidation of Metaxylol-sulphamids.—Oscar Jacobsen.—A controversial paper with reference to the communication by Iles and Remsen (*Berichte*, xi., 1326).

Dibrom-metaxylol-sulphonic Acid.—O. Jacobsen and E. Weinberg.—The authors examine the soda-salt, the acid chloride, and the amid.

On Paraxylidin.—W. Schaumann.—Paraxylidin is an oily liquid lighter than water, colourless when recently prepared, but turning yellow on exposure to the air; abundantly soluble in hot water, and boiling at 220° to 221° . It forms with acids well-defined salts.

Action of Potassa upon Tetra-nitro-diphenyl-urea.—S. M. Losanitch.—The result of the reaction is a green, solid body of the composition $C_{13}H_6N_6O_9K_2$.

On Lotura Bark.—O. Hesse.—The author has isolated from this bark three alkaloids, which he names loturin, colloturin, and loturidin, the properties and compounds of which he here describes.

Certain Substitutes for Quinin.—O. Hesse.—The Australian tree *Alstonia constricta* was formerly supposed to contain quinin; recent observation has instead merely detected alstonin, a bitter principle not possessing basic properties. Ditain, a principle obtained from the bark of *A. scholaris* is also no alkaloid. Crossopteryx bark has been stated to contain quinin, but in the author's opinion it is little better than bad fire-wood.

Remarks on Mr. Rice's Memoir on the Cinchona Alkaloids.—O. Hesse.—The salts of quininid, which Rice takes to be conchinin, do not contain the latter base.

Reduction of Aceto-phenon.—K. Buchka.—A preliminary reply to Engler's paper (*Berichte*, xi., p. 934).

Adulteration of Wine.—F. v. Lepel.—The author gives the spectroscopic reactions of a mixture of magenta and the juice of *Beta vulgaris*.

On Aurin.—R. S. Dale and C. Schorlemmer.—The authors explain the formation of aurin by the following equation:—



Determination of Nitric Acid as Ammonia.—E. A. Grete.—The author, on heating pure saltpetre with xanthogenate and soda-lime, obtained almost exactly the whole of the nitrogen present in the state of ammonia.

MISCELLANEOUS.

Royal Institution of Great Britain.—The Managers have decided that the next Actonian Prize shall be awarded in 1879 to an Essay illustrative of the Wisdom and Benevolence of the Almighty; the subject being "The Structure and Functions of the Retina in all Classes of Animals, viewed in relation with the Theory of Evolution." The Prize is One Hundred Guineas, and will be awarded or withheld as the Managers shall think proper. Competitors for the Prize are requested to send their essays (with or without their names being affixed) to the Royal Institution, addressed to the Secretary, on or before October 1, 1879. The adjudication will be made by the Managers in 1879.

NOTES AND QUERIES.

Oils.—(Reply to R. J.).—I think that your correspondent R. J. will find the information he requires in the article on Oils in "Ure's Dictionary." F. C. Calvert has also published a paper in the Pharmaceutical Society's *Journal*, xiii., 56, upon the "Analysis and Detection of Impurities in Oils."—H. A. LAWRENCE.

TO CORRESPONDENTS.

Max Jordan.—We regret that we are unable to give our correspondent the information he desires.

Lavon.—Morfit's treatise on Soap, published by Trübner and Co.

Merrick.—Apply to the Secretary of the Society at Burlington House for a nomination form, which has to be signed by a certain number of Fellows of the Society, some of whom must have personal knowledge and others general knowledge of a candidate.

J. C. Harvey.—"Sutton's Volumetric Analysis," published by Churchill.

THE CHEMICAL NEWS.

VOL. XXXIX. No. 1000.

SILICIURETTED HYDROGEN.

We have received from Dr. Theodor Schuchardt, of Goerlitz, a specimen of a new body, which he calls silicium strontium. It is formed from the preparation of metallic strontium by electrolysis, but no particulars are given as to the substances present or the reaction by which it is formed. As received from Dr. Schuchardt the compound is a grey powder with a slight odour resembling phosphuretted hydrogen. When mixed with dilute hydrochloric acid a rapid evolution of the spontaneously inflammable siliciuretted hydrogen takes place. No particulars as to price are mentioned, but if obtainable in quantity this compound will probably be the readiest source of siliciuretted hydrogen.

W. C.

MEANS OF DETECTING THE ADULTERATION OF SAFFRON.

To detect adulteration with *Calendula flowers* (Feminelle), it is merely required to moisten a few flowers, and to rub them singly with the finger on white paper. The genuine flowers will give a fine rich yellow colour, whilst the Feminelle will only yield a violet reddish hue. It can also be easily detected by soaking the suspicious flowers in pure or, better still, distilled water. The real saffron will retain its fine red colour after hours, whilst the Feminelle will lose its artificial tint within a short time.

To detect an admixture of *Honey and Barytes* it is merely required to put a pinch of saffron in a tumbler with pure clear water (also in this case distilled water is preferable), and agitating it for a few minutes. Adulterated saffron will at once turn the water cloudy, and even small particles of dust may be seen falling to the bottom, which, on pouring the water carefully out, will be found to be a slimy, sand-like mass. With pure saffron the water will remain clear, showing a fine pure yellow colour, which, according to the quality of the flowers, will be more or less intense. Five or ten minutes suffice for these experiments.

PRELIMINARY NOTE ON THE ACTION OF SEA-WATER ON THIN IRON AND STEEL PLATES.

By SERGIUS KERN, M.E., St. Petersburg.

Thin iron and steel plates were subjected to the action of sea-water in the following manner:—

The specimens were 0.076 metre square and 0.00317 metre thick (one-eighth of an inch). They were placed (four samples of iron, four samples of steel plates) in wooden boxes containing sea-water, 2 litres in each box. The sea-salt was obtained from the north shores of France and dissolved in water; the specific gravity was 1.027 when the salt was added.

The specimens were immersed in this solution for thirty days, next dried between filter paper, and finally in an air-bath at 35° to 40°; then they were weighed.

All the plates before the experiments had a smooth surface, and were of the best manufacture. The chemical composition of them being as follows:—

I. Steel Plates.

	Per cent.			
	I.	II.	III.	IV.
Carbon ..	0.290	0.300	0.180	0.234
Manganese ..	0.380	0.412	0.275	0.178
Sulphur ..	0.010	0.010	0.012	traces
Phosphorus ..	0.005	0.008	0.010	0.032
Silicon ..	0.015	0.011	0.010	0.021

II. Iron Plates (Average Analysis).

Carbon ..	0.110 per cent.
Manganese ..	0.187 "
Sulphur ..	0.010 "
Phosphorus ..	0.004 "
Silicon ..	0.008 "
Copper ..	traces "

In the steel plates, in Nos. II. and III., traces of copper were observed. The other two steel plates contained no copper.

In the annexed table is shown the loss of the plates during the action of the sea-water upon them.

III. Action of 100 litres of Sea-water (Sp. Gr. 1.027) on one Square Metre of Iron and Steel.

1-inch Plates.	Loss in Grms.
1. Steel ..	22.75
2. " ..	22.34
3. " ..	21.83
4. " ..	21.96
1. Iron ..	25.78
2. " ..	25.54
3. " ..	26.04
4. " ..	27.44

To come to some conclusion further experiments are wanted, which are at the present time commenced, but the author believes that it is preferable to use steel for the skin of the ships. Certainly, in using steel much will depend on the manufacture and the composition of it, and many equivocal results may be obtained during experiments on the action of sea-water, owing to the more or less compactness of the plates. The best control in this case may be the determination of the specific gravity of the plates.

ON THE DIRECT PROCESS OF MAKING WROUGHT-IRON AND STEEL.*

By CHARLES M. DU PUY, C.E.

ABOUT one year ago I had the honour of addressing this Institute on the direct manufacture of iron from the ore. I then briefly referred to the various efforts that had been made in that direction, reaching over the best part of the past century, and closed the paper by describing a method, which, from various experimental and practical tests, seemed likely to promise usefulness in the arts.

It may be remembered, by this process the ore, carbonaceous matter, and fluxes, in the proper proportions, are ground together and mixed at the same time, and then filled into annular sheet-iron cases, holding from 100 to 200 pounds. Ten to twenty of these cases, with spaces between them, are subjected to the gradually increasing heat of a reverberatory furnace, and in about five hours on the average the ore and cases settle down, without work upon them, to about one-third their original height, becoming welded into quite compact lumps of iron, interspersed with liquid slag. These metal lumps are either then removed separately from the furnace, or several of them being welded together at the same heat, are forged or squeezed and rolled to "muck-bar."

* A Paper read before the Franklin Institute, November 20, 1878.

This process differs from all others, and it is exactly the reverse of the forge-fire and the blast-furnace methods. In those, a tuyere conducts a stream of oxygen into close contact with the particles of ore, while in this process the moderate heat prevents the combination of phosphorus with the iron, while atmospheric contact is excluded during reduction, and a reflected heat from the roof is furnished, for absorption during the transition to metal. It is to this peculiar treatment that is ascribed the high quality of the iron for steel purposes. It is what may be termed a *baking* process, which is conducted at a comparatively moderate heat to *eliminate phosphorus*, and in which the ore is kept out of the reach of oxidising influences, and to this the high value for steel is attributable.

At the time I addressed you I had licensed Messrs. Miller, Metcalf, and Parkin, of Pittsburg, Pa., to work the process at their Crescent Steel Works. Their forge and furnace was not completed until some time in last January, when for several months thereafter they carefully tested the iron which was deoxidised with charcoal by this method, in the various ways customary to determine the value of iron for high grades of steel. The result of these investigations brought forth the unqualified endorsement of this firm that the iron so produced from our native ores, for steel purposes, is equal in every respect to the most costly grades of Swedish iron.

Although the production of fine steel iron, reduced with charcoal by this process, was a step forward, yet the consumption of high grades of steel is comparatively limited, being mostly used only for tools of various kinds requiring great endurance. In order to enlarge the uses of this superior make of iron, it became apparent that its cost must be cheapened by deoxidising the ore with waste anthracite dust or refuse coke-dust, instead of charcoal.

To determine the practical value of anthracite dust, a series of more than fifty experiments were conducted, during the month of August last, at the forge at Reading, Pa. In these experiments, with anthracite dust, were used magnetic ore from near West Point, N.Y., Dickinson ore from northern New Jersey, Cornwall ore from Lebanon Co., Pa., Cumberland Valley ore from Maryland, Hematite ore from near Newark, Del., besides several other ores found near the line of the Philadelphia and Reading R.R.

These ores were worked both separately and variously combined, and with one exception, in every case, were reduced and forged to blooms. The blooms, which were re-heated, were drawn out smoothly under the hammer. The tensile strength of one of these bars was tried, and found to not exceed 45,000 pounds to the square inch.

A couple of crucibles of steel were made from a part of the iron which had been re-heated and drawn out, and the ingots were forged and used for planing tools. They stood forging and tempering well, comparing favourably, in endurance, with steel usually used in planing iron.

The object of testing iron reduced from the ore with *anthracite* coal-dust, for tool steel, was not so much with the expectation that at the first trial it would work well in planing iron in competition with steel carefully prepared from *charcoal* iron, but to determine, in a general way, the quality of the metal for steel purposes, of superior quality and in large quantities, by the open hearth. The result of this crucible test surpassed expectations and was very gratifying, from the fact that it is believed to have been the first instance where good tool steel was ever made by a direct process, on a practical scale, from iron deoxidised with *anthracite* coal-dust, and confirms the belief that anthracite dust will eventually be largely used to produce steel of high quality, cheaply and in large quantities.

Besides testing the ores before named, at the Reading forge, iron scale from the rolls (which is almost pure oxide of iron) was reduced alone with anthracite dust, and was also mingled with ore and anthracite, and in both cases was found to forge well into good blooms.

At the same time with the foregoing experiments, the ul pho oxides or refuse ore remaining after extracting

sulphur for sulphuric acid from the iron pyrites, at the New Jersey Chemical Works (commonly known to the trade as "Blue Billy"), was also reduced by this process with anthracite coal dust, and forged well into a bloom. This specimen was some time afterwards re-heated, rolled, cut up, and piled with about one-third of its weight of muck-bar from common puddled iron, and plated out well into smooth sheets of No. 26 iron. This last experiment is not of much value for this country, because very little sulphuric acid is made from pyrites, but in Europe probably not less than a million tons of this refuse ore is now annually almost wasted, because of the difficulty of economically utilising it.

These Reading experiments were the more gratifying from the fact that the process was quite successfully conducted in a reverberatory furnace, not well adapted to the purpose. This furnace had some time before been specially arranged to test the use of anthracite coal-slack, by burning it in fires of 4 or 5 inches thickness, on what is commonly termed "the Wooten grate-bars." As these thin fires required renewal every 15 or 20 minutes, the frequent opening of the door for replenishment exposed the ore so often to atmospheric re-oxidation as to make the furnace unsuited to the process. Besides this, the fire and flue bridge-walls, which had been constructed very low, and could not be very well raised owing to the lowness of the roof, caused the cases of ore to be exposed to the direct action of the blast and draught as it passed through the furnace into the stack.

Following up these interesting experiments during the month of September last, a sand-bottom scrap-heating furnace was altered at the Sligo Iron Works at Pittsburg for the purpose of testing this process further, by throwing the lumps of metal into a Burden Squeezer, and then rolling them to "muck-bar" at the same heat. Hitherto they had usually been forged to blooms under a hammer, and afterwards re-heated to be drawn to bars.

This furnace was operated experimentally with 32 heats. There was found no difficulty in making balls to pass through the squeezer and muck rolls at the same heat, just as ordinary puddle balls from pig-iron is squeezed and rolled at the same heat to muck-bar; but in order to fill the squeezer, which required balls of 150 to 200 pounds to secure a good compression, it became necessary to weld and press several of the lumps of metal together in the furnace at a heat so high as to cause the alkali to drip more or less from the mass, and soften the sand-bottom. As the balls were thus compressed and rolled in the fused sand an unnecessarily large portion of the iron was thereby cut by the sand and wasted to a silicate of iron, so that the yield from Republic ore which, before the bottom became softened, produced 53 pounds of iron in muck-bar from 100 pounds of ore charged, and which it was expected would have been brought up to 60 pounds of iron from 100 pounds of ore, gradually became very much lessened, showing conclusively that reduced yield followed the softening of the bottom, and determined the necessity of a "cinder-bottom" in order to save the iron.

To compress and weld several of these masses of metal together in the furnace, in order to have them enter and fill the squeezer, *really comprised all the labour required during the heat*. The cases were 15 inches diameter and 14 inches high, holding about 135 pounds of ore besides the coke and fluxes. By making the cases very little larger, say 16 inches diameter and 16 inches high, they would each produce about 100 pounds of iron, and by withdrawing each separately to the squeezer, all furnace manipulation of the metal would be avoided. Then, aside from maintaining the fire and charging and discharging the metal, all the labour really needed during the heat would be to change the damper once or twice, enabling an ordinary heater to regulate all the furnaces in the largest mill.

The fire- or grate-surface of this Pittsburg furnace was 4 feet square, and the space between the bridge-walls was 9 feet by 5 feet in width. The bottom was of sand,

set hard by heat. The flue-bridge was built up within 3 inches of the roof. The stack was 18 inches square inside. A fan-blast was used, part of the air passing under the grate-bars, and part, being heated, was passed through and over the fire-bridge.

In these several experiments it was found that, although the opening between the flue-bridge and roof for the escape of the spent products of combustion into the stack was lessened more than one-half the size from what it was when, with less capacity, the furnace had been used for heating scrap-iron, still the heat was so intense as to soften and render useless the damper rod at the top of the stack, which had previously stood twelve years, requiring renewal three times during these experiments. The conclusion drawn from these accidents was that the grate-surface of this furnace was amply sufficient to supply the accumulating heat for a length of 14 or 15 feet between the bridge-walls instead of 9 feet, thereby nearly doubling the capacity of the furnace with the same consumption of fuel.

Expensive fluxes, such as soda, manganese, &c., have been abandoned, as it has been now conclusively proved that a proper mixture of aluminous and silicious ores with lime, to produce a non-flowing glassy slag, is all that is required. This glassy slag being mingled among the particles of ore, covers and seals them from re-oxidation from furnace heat during the process of reduction. When either of these glass-making constituents are not available in the proper proportion, in the ore, ordinary sand, or clay, in the proper proportions, may be added to the lime and mixed together without injury to the metal, as the glass is readily compressed out of the iron by the hammer or squeezer.

During these experiments No. 26 sheet-iron was rolled out at the mill from common puddle bar, being found to answer equally as well for cases as well finished sheets from the best iron.

It has now been demonstrated that good steel can be made by this process from iron deoxidised either with charcoal, anthracite-dust, or coke-dust. The carbon in all these fuels being alike, and as the ash of either, along with the earthy impurities of the ore, be they more or less, may be readily reduced to a glass by a proper mixture of lime and ores through chemical analysis, and so utilised as a covering of the metal during deoxidation, and compressed from it afterwards—it follows the truest commercial economy is found in using that form of carbon which is of least value.

In the coke-making bituminous regions, the refuse coke-dust is disposed of in waste heaps and often burnt; while from the beginning of anthracite mining, mountains of anthracite slack remain standing monuments of our wastefulness. These wasted fuels by this process offer a premium for the production of cheap steel.

Any process that will not withstand the closest economic scrutiny is justly recognised as valueless in the arts; it is therefore proper to remark that the investigations at Reading and Pittsburg during the past year, although conducted in furnaces not specially adapted to the process and often amid inconveniences inseparable from experiments surrounded by the activity of regular mill-work, still nevertheless substantially confirm the view heretofore asserted, that a true commercial economy cannot fail to result from a systematic working of the process.

When it is considered that this mode of making iron frees it almost entirely from phosphorus, which goes almost altogether into the slag as explained in my former paper.

That a furnace at present prices for material and labour, in which from 1 ton to 1½ tons of blooms may be produced every twenty-four hours, will not cost over 800 dollars to 1000 dollars.

That a little lime in combination with ore and waste coal-slack has been substituted for more expensive fluxes and is found sufficient to supply the glassy covering for the metal.

That No. 26 sheet-iron for cases is now found to be

good enough when plated out from common puddle bar, and that the cases are very simply and cheaply made, needing neither tops nor bottoms.

That two men, in ten hours, with a 15 horse-power engine, will grind, mix, and fill the cases with the ore mixture for 7 tons of blooms.

That the furnace work is confined to charging, discharging, and firing the furnace, there being no necessity whatever for manipulating the metal from the time it is charged in the furnace until brought to the hammer or squeezer, thus greatly simplifying and cheapening what by the puddling process is excessively laborious and expensive work.

That the masses of metal, as one by one they are squeezed to blooms, may then be transferred in the heated state to the open-hearth bath, and there quickly melted along with the other steel-making material for the highest quality of steel.

When all these points are well considered it may be clearly understood that in localities as favourably situated for instance as the line of the Philadelphia and Reading Railroad, where Cornwall and other ores can be laid down at an average of 3 dols. per ton, and where anthracite fuel and coal-dust may be obtained proportionally low—that blooms on a large practical working scale, nearly freed from phosphorus, may be manufactured equally as good for 18 dols. or 20 dols. per ton, as now cost from 38 dols. to 50 dols. per ton. Similarly great inducements are likewise offered in Northern New Jersey, where fuel and ore can be brought together at about the same cost. In the bituminous coal region fuel is correspondingly low, and coke-dust can be had for the freight, offering favourable manufacturing advantages.

In a few words, this process offers a cheap mode of making a high quality of metal for steel, by an inexpensive plant, enabling manufacturers to utilise idle rolling mills at trifling cost.

THE SEVEN FUNDAMENTAL TYPES OF ORGANIC CHEMISTRY,

AS VIEWED AND INTERPRETED FROM THE STANDPOINT OF THE "TYPO-NUCLEUS" THEORY.

By OTTO RICHTER, Ph.D.

IMPRESSED with the deep scientific import and significance of certain far-reaching and well-matured original ideas and speculations of my own, I have deemed it my duty of again addressing the Chemical Profession in order to submit to them for closer examination a second and even more abundant crop of valuable and interesting generalisations which I have been able to gather on the wide and fruitful field of speculative research. It may not be amiss to state here that the task set before me, and commenced in full earnest some thirty years ago, was persevered in with the avowed object of contributing my share also towards the foundation of a more rational, comprehensive, and harmoniously elaborated system of chemical philosophy. As a further proof of the sincerity of my motives and aspirations, I have now, I flatter myself, succeeded in concocting an entirely novel and original theory of molecular substitution, which promises to meet all the more pressing wants and requirements of modern organic and inorganic chemistry. It is with the aid of this latest and still vigorously expanding offspring of my private meditations, reared as it has been under the fostering care of those purer doctrines and principles which I had occasion to broach and to discuss in a series of papers under the head of my "Typo-Nucleus" Theory, that I hope to shed a bright and powerful light upon sundry vexed and seemingly insoluble questions, concerning which the

History of our science has recorded so many stirring, but in their final results comparatively barren and unprofitable, controversies. Having, in the course of my speculative travels, made the startling discovery that a grave and all-pervading error had been committed on the part of our leading theorists in not sufficiently discriminating between two in their *modus operandi* essentially different chemical processes, I have considered it imperative on me to expose this fatal error, the offshoot no doubt of an overbold and over-hasty generalisation, and, as far as lay in my power, to counteract its evil influences and tendencies.

The two processes just alluded to, and which are constantly encountered in our laboratories, and not unfrequently in one and the same reaction, are on the one hand the process of direct chemical union by the addition of new elements to those already combined; and, on the other hand, the process of indirect chemical union by the substitution of new elements for those already combined. Now, I venture to maintain, that our chief chemical authorities and advisers, from neglecting to draw a clear and distinct line of demarcation between these two radically different methods of proceeding, have beguiled themselves and their numerous followers into the adoption of a style of writing which, by inundating the chemical market with a seemingly very plausible but really unsound and unpalatable species of sham literature, threatens not only to create a distaste for more solid work, but likewise to seriously retard the healthy growth and development of theoretical chemistry.

With these well-intentioned and, I am afraid, well-founded strictures and objections, I shall at once proceed to enlarge on some of the more salient doctrines and principles of my own system, a brief exposition of which will, I trust, enable the reader to appreciate the true merits and scientific value of my new substitution theory.

In striking contrast with the prevailing biatomic hypothesis, according to which the different kinds of elementary molecules are represented as consisting of two chemically identical atoms, cemented together by means of one, two, three, or more reciprocally attracting centres of force, my system is founded upon the polyatomic hypothesis, according to which the different kinds of elementary molecules, though likewise regarded as being composed of an equal number of atoms, are at the same time believed to enclose within them a considerably larger number of such atoms than are generally assumed on the other side. These primordial, and in all probability spherical, particles of ponderable matter are held to differ from each other in point of size, specific gravity, and electro-polar energy, according to the species they belong to, but they have, each and all, this highly characteristic feature in common, that they are symmetrically disposed and grouped together in their respective molecules with reference to the three conjugate axes of space, and in obedience to some omnipotent fundamental law of atomic collocation and arrangement. Another remarkable and noteworthy feature of the molecules before us, which in my notation are expressed by the general symbol E_2 consists in this, that, while they are not held to correspond to entities enjoying a separate and independent state of existence, they are at the same time held to be of immense theoretical value and importance, inasmuch as they furnish the only solid and substantial basis on which I believe it possible to erect a truly rational and comprehensive system of chemical philosophy. A select number of these theoretically indispensable molecules have, as a reference, been embodied in the accompanying table, together with their respective symbols and atomic weight-equivalents, and it deserves special mention that the particular order in which these symbols are made to succeed each other is intended to convey an approximate idea of the relative electro-polar positions which the corresponding molecules are believed to occupy in the atomic edifice in conformity with the general electro-polar law of molecular collocation and arrangement.

Table of Chemical Elements, with their Symbols and Atomic Weight Equivalents.

First Division: Electro-positive Elements or Metals.

Symbols . . .	H ₂	K ₂	Na ₂	Ba ₂	Ca ₂	Mg ₂	Zn ₂	Pb ₂	Fe ₂	Cr ₂	Al
Atomic weights	2	78	46	137	40	24	65.5	207	56	52	27.5
Symbols . . .	Ag ₂	Pt ₂	Au ₂	Sn ₂	Hg ₂	Bi ₂	Sb ₂	As ₂	P ₂	—	—
Atomic weights	216	197.5	197	118	400	210	122	75	31	—	—

Second Division: Electro-negative Elements or Non-metals.

Symbols . . .	Si ₂	C ₂	Bo ₂	Te ₂	Se ₂	S ₂	T ₂	Br ₂	Cl ₂	Fl ₂	N ₂	O ₂
Atomic weights	28	12	11	128	79.5	32	254	160	71	38	14	16

In commenting on the contents of the preceding table I am far from maintaining that the relative places assigned to the various elements in the general electro-polar order of grouping are absolutely correct. I can only express my regret that, for want of proper experimental data, my researches in this direction have not as yet led to any satisfactory results, and I am bound to confess that, in order to fill up the numerous gaps, I have reluctantly allowed myself to be guided by the uncertain glimmer of fancied analogies and probabilities. Thus, for instance, I am unable to answer the question why K₂ should occupy the second place in the electro-polar order of grouping in preference to the third or fourth place; and, similarly, why Zn₂ should precede Pb₂ or any other metal. But, on the other hand, I hope to prove, with the aid of a great variety of valuable experimental evidence, that H₂ is really the most electro-positive of all the known elements, whereby it claims for itself the first or lowest place in the electro-polar order of grouping, while O₂ is really the most electro-negative of all the known elements, whereby it claims for itself the last or highest place in that order. I am further able to prove that the members of the non-metallic series—C₂; S₂; I₂; Br₂; Cl₂; Fl₂; N₂; O₂—are given in their proper order; while I think it highly probable that the members of the metallic series—Bi₂; Sb₂; As₂; P₂—are really succeeding each other in the manner indicated. This conjecture derives considerable support, at all events, from the unmistakable analogy which the series before us bears to the series I₂; Br₂; Cl₂; Fl₂, the atomic weights of whose members are observed to diminish as we ascend in the electro-polar order of grouping, a coincidence which, if not accidental, must be the outcome of some general law, to the jurisdiction of which not only the series in question, but various other groups of cognate elements, may likewise be found amenable.

Trusting that this particular field of speculative research may soon become better known and cultivated, I shall now proceed to describe the general method whereby that class of elementary molecules which form the theoretical basis of reasoning may become transformed into that other class of elementary molecules which are capable of enjoying a separate and independent state of existence, and through them into that endless multiplicity and diversity of compound molecules, the internal structure and organisation of which still continues to defy the argumentative skill and ingenuity of our shrewdest and most keen-sighted speculative chemists and physicists.

In pondering this momentous and seemingly inaccessible problem my efforts were at length rewarded with the important discovery that directly or indirectly the former class of molecules are destined to furnish the materials for the construction of seven distinct fundamental types or forms of molecular grouping, which, by their combination with each other, are further destined to furnish a vast number and variety of composite types, under which the whole immensity of organic and inorganic compounds may be systematically arranged and classified. The proper treatment of this subject requires that I should divide the before-mentioned seven fundamental types into three separate sets on the ground that each set owes its origin and formation to the aid and intervention of a distinct fundamental principle or agency. Accordingly, the first set of types is believed to stand under the immediate control of the principle of parity or physical principle, and

to embrace three fundamental types called respectively the Nucleus Type, the Outer Conjunct Type, and the Inner Conjunct Type. Again, the second set is believed to stand under the immediate control of the principle of calority of thermal principle, and to embrace two fundamental types, called respectively the Outer Subjunct Type and the Inner Subjunct Type. Finally, the third set is believed to stand under the immediate control of the principle of polarity or chemical principle, and to embrace two fundamental types, called respectively the Outer Adjunct Type and the Inner Adjunct Type.

(To be continued)

PROCEEDINGS OF SOCIETIES.

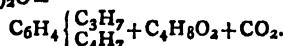
CHEMICAL SOCIETY.

Thursday, January 16, 1879.

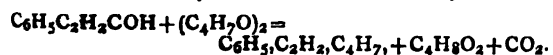
Dr. J. H. GLADSTONE, F.R.S., President, in the Chair.

AFTER the announcement of visitors, the minutes of the previous meeting were confirmed. The following certificates were read for the first time:—W. Spottiswoode, P.R.S., T. Whitaker, J. L. MacMillan, J. A. Ogilvie, W. S. Lawson, V. H. Veley.

The first communication was made by W. H. PERKIN, F.R.S., "On the Action of Isobutyric Anhydride on the Aromatic Aldehyds." When an aromatic aldehyd is heated with normal butyric anhydride and a butyrate an angelic acid is obtained containing the hydrocarbon radical of the aldehyd employed. Thus, benzoic aldehyd yields phenyl-angelic acid, &c. The author has studied in the present paper the reaction which takes place when isobutyric is substituted for butyric anhydride. A mixture of cuminic aldehyd, isobutyric anhydride, and sodium isobutyrate was heated in sealed tubes to about 150° C. for twenty-four hours. On opening the tubes much carbonic anhydride escaped. The product was mixed with water, and boiled in a retort until the distillate was nearly free from oily matter. On cooling, the aqueous portion was separated from the thick oily product, and the latter boiled with an excess of a solution of sodium carbonate. The boiling alkaline solution was acidified, and deposited a thick oily acid. This was dissolved in petroleum spirit, and after some time the solution deposited oblique crystals. These proved on analysis to be a compound of cumenyl-crotonic and isobutyric acids. A similar experiment with benzoic aldehyd gave phenyl-crotonic acid. When isobutyric acid, prepared by repeated fractioning, and so, as far as possible, freed from propionic anhydride, was used, very much smaller quantities of the crotonic acids were obtained. The author estimated the amount of carbonic anhydride produced in the above reaction, and examined the oily matter which distilled over on boiling the products of the above reactions with water. It proved to consist principally of butenyl benzene, $C_{10}H_{12}$ (when hydride of benzoyl was used), boiling at 184° to 186°, yielding, on oxidation with chromic acid, benzoic and acetic acids. The author proposes to call it β -butenyl-benzene. Its dibromide and bromo derivative and the dibromide of the latter were prepared and examined. With cuminic aldehyd β -isopropyl-butenyl-benzene was obtained,—



It boils 8° lower than the α body. When cinnamic aldehyd was used, butenyl-cinnamene was obtained,—



It oxidises rapidly, and forms a red crystalline compound with picric acid. By heating the sodium derivative of or the hydride of salicyl itself, with isobutyric acid, ortho-hutenyl-phenol, $C_{10}H_{12}O$, was obtained as a colourless oil, boiling 223° to 225°, having a smoky and cedar-like odour; specific gravity = 1.0171. Many reactions of this substance are given in the paper. By treating paroxy-benzoic aldehyd in a similar way para-butenyl-phenol was obtained, boiling at 230° to 235°, crystallising at low temperatures. Similarly, from anisic aldehyd, β -para-butylen-anisole was obtained, fusing at +7°, $C_{11}H_{14}O$; it boils at 236° to 237°. In conclusion the author discusses the isomerism existing between the bodies obtained in the present research and those obtained previously by using normal butyric acid, and especially that of the butenyl benzenes.

After some remarks by the PRESIDENT and Drs. FRANKLAND and ARMSTRONG on the theoretical part of the above paper,

Dr. DUPRÉ read a communication "On Two New Methods for the Estimation of Minute Quantities of Carbon: (1) Gravimetric; (2) Chromometric; and their Application to Water Analysis," by A. DUPRÉ and H. WILSON HAKE. (1) The Gravimetric Method:—This method consists essentially in burning the small quantity of carbon in a stream of oxygen in an ordinary combustion-tube containing some granulated cupric oxide. One end of the combustion-tube is drawn out and bent downwards at an angle of 120°, so that it can easily be attached to a Pettenkofer tube charged with a perfectly bright solution of barium hydrate. The stream of oxygen passes through a long tube containing caustic potash before entering the combustion-tube, and before commencing the combustion the tube with the cupric oxide is heated to redness, and the oxygen passed until carbonic acid ceased to be evolved, as tested by passing through bright baryta-water. The Pettenkofer tube is charged with baryta-water, various precautions being taken to prevent the slightest access of any air containing carbonic anhydride, and connected with the combustion-tube. The substance is introduced and burned in the usual way. The carbonic acid is completely absorbed by the baryta-water. At the conclusion of the combustion the barium carbonate formed is filtered and washed by an ingenious arrangement completely out of contact with ordinary air, first with water saturated with barium carbonate, and finally with pure water. The Pettenkofer tube is then rinsed out with dilute hydrochloric acid, and the washings poured on the filter containing the barium carbonate. The barium chloride solution is then evaporated in a very small platinum dish, converted into sulphate by ignition with sulphuric acid, moistened with nitric acid, again ignited, and weighed on an assay balance. The following are results thus obtained with known quantities of sugar:—

Carbon Taken.	Carbon Found.
0.01344	0.01340
0.00755	0.00748
0.00535	0.00551
0.00390	0.00398
0.00137	0.00163
0.00079	0.00085
0.00034	0.00027

The method is therefore quite accurate enough for determining the carbon obtained from a litre of a first class potable water. The authors found the error of several blank experiments to be equal to three-tenths of a milligramme of carbon: they therefore subtract this quantity from each result. The chromometric method, or, as the authors name it, the nephelometric method, consists essentially in burning the carbon as above, but the carbonic acid is conducted into a standard solution of basic acetate of lead (2 per cent), and estimating the turbidity produced, as compared with that produced by the carbonic acid evolved under similar circumstances by a known and nearly equal quantity of carbon (sugar), the difference between the two being estimated by a Mills's colorimeter.

The authors find that the precipitate produced in the acetate of lead is very similar in appearance under widely different circumstances, and that it settles slowly. This method is extremely delicate, and a difference produced by 3-100ths of a milligram can be clearly estimated. The authors also propose, to prevent any chance of loss in scraping out a residue, the use of a collapsible silver-foil dish, which is placed inside a platinum dish for support, and when the residue is dry the silver dish with the residue is rolled up, and the roll introduced bodily into the combustion-tube. These dishes hold about 50 c.c. (They are made by Johnson and Matthey, and cost about 3s. 6d.) Using these methods, the authors estimated the carbon in a water, using 200, 100, 50 c.c., with the following results:—

Carbon in residue from 200 c.c. water gravimetric method equal 0.34 per 100,000.

Carbon in residue from 100 c.c. water nephelometric method equal 0.31 per 100,000.

Carbon in residue from 50 c.c. water nephelometric method equal 0.26 per 100,000.

It must be understood that the authors do not recommend the use of such small quantities of water, but the results show the delicacy of the processes. In conclusion, the authors suggest the use of the above methods for estimating the carbon in iron and the carbonic acid in air.

Dr. FRANKLAND said that every chemist must feel indebted to the authors for the very ingenious methods they had brought forward, methods which seemed to be sufficiently accurate and delicate for the purpose intended: there still remained, however, the estimation of the nitrogen, which was of equal importance with the determination of the carbon. He would like to ask whether the condition of the precipitate in the plumbic acetate did not vary somewhat with different conditions, *e.g.*, the rate at which the carbonic anhydride was evolved.

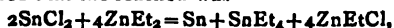
Dr. ARMSTRONG asked how the complete combustion of all the carbon could be ensured with the silver dish? Alkaline carbonates might be formed.

Mr. THORPE said that he had scraped out many water residues, and had never experienced any difficulty in the operation. An apparatus similar to that used by the authors had been employed by Fleming at Cheltenham College. He, however, decomposed the barium carbonate, and read off the carbonic anhydride evolved.

Prof. HARTLEY remarked that Dittmar had suggested methods for easily estimating the C and N in water residues: he absorbed the carbonic acid by soda-lime. The nitrogen was converted into ammonia by soda-lime, the ammonia absorbed in weak acid and Nesslerised.

Dr. DUPRE replied that the precipitate in the basic acetate of lead seemed to be remarkably uniform; that the water was evaporated with phosphoric acid, so that no alkaline carbonates could be formed; that although the residue from a litre of water might easily be removed, that from 50 c.c. could hardly be detached and transferred without loss.

Dr. FRANKLAND then read a communication "*On Stannic Ethide*," by E. FRANKLAND and A. LAWRENCE. In endeavouring to prepare stannous ethide by the action of zinc ethyl upon dry stannous chloride the authors discovered that the reaction was—



and that by its means stannic ethide could be prepared more conveniently and in larger quantities than has been possible hitherto. Fragments of fused stannous chloride were added to zinc ethyl contained in a flask cooled by immersion in water. As soon as the mixture ceases to fume in the air, when a sample is taken on the end of a glass rod, enough chloride has been added. The pasty mass is distilled in an oil bath. The distillate, which should contain some free zinc ethyl, is poured into water, the zincic hydrate dissolved in dilute sulphuric acid, and the heavy oily layer of crude stannic ethide separated and purified. The pure substance has, at 180° C., no action on aluminium, sodium, magnesium,

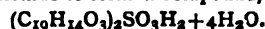
acetone and ethylic oxalate. It acts slightly on benzoic aldehyd. At ordinary temperature neither ammonia, carbonic anhydride, carbonic oxide, cyanogen, nitric oxide, oxygen, nor sulphuretted hydrogen affect it. Sulphurous acid is slowly absorbed, a crop of crystals being formed after some weeks. These were soluble in ether, and proved to be stantriethyl sulphate, identical with the sulphate of sesquistanethyl of Cahours and Buckston. The syrupy liquid, from which the above crystals had separated was found to contain stantriethyl hydrate combined with ethyl-sulphonic acid. The reaction was probably— $\text{SnEt}_4 + \text{SO}_2 + \text{O} = \text{SO}_2\text{Et}(\text{SnEt}_3\text{O})$. The reaction was tried when air was rigorously excluded during the absorption of SO_2 ; but the final products were the same; much metallic tin was, however, formed. The authors therefore infer that oxidation occurred during the solution and evaporation of the products.

The PRESIDENT said the above paper furnished us with a ready method of preparing stannic ethide, so that we could obtain it with facility if it was required as a reagent; it was singular that it should resist the action of so many substances.

The next paper was read by the SECRETARY; it was entitled "*On Aurin*," Part II., by R. S. DALE and C. SCHORLEMMER. In previous researches the authors stated that by the action of alcoholic ammonia on aurin, rosanilin was obtained, and a dilemma arose, that if aurin had the formula $\text{C}_{20}\text{H}_{14}\text{O}_3$, rosanilin could not have the formula which Hofmann had proved it to have, $\text{C}_{20}\text{H}_{19}\text{N}_3$. The authors therefore again prepared pure aurin and confirmed their previous analysis. Gräbe and Caro also assigned the same formula to aurin. Zulkowsky states, however, that the method employed by the authors yields wretched results, and cannot be used on the manufacturing scale. The authors, however, affirm in the present paper that under certain conditions, which they are not at liberty to divulge, a fairly good yield can be obtained. The authors have used two methods to purify aurin; one by conversion into ammonia aurin and the decomposition of the latter by hydrochloric acid, and a second by taking advantage of the fact that the solubility of aurin in alcohol decreases with the removal of the by-products. The analyses of pure aurin thus obtained completely confirmed the formula $\text{C}_{20}\text{H}_{14}\text{O}_3$. The correctness of the formula was also proved by E. and O. Fischer, who explained the above discrepancy by showing that the rosanilin obtained from aurin was para-rosanilin, $\text{C}_{19}\text{H}_{17}\text{N}_3$. The authors next consider the formation of aurin, which they consider takes place as follows:—



The paper contains the examination of several compounds and derivatives of aurin. Ammonia aurin, $\text{C}_{19}\text{H}_{14}\text{O}_3(\text{NH}_3)_2$. Tetra-brom-aurin, $\text{C}_{19}\text{H}_{10}\text{Br}_4\text{O}_3$. The authors have studied the action of acetyl-chloride and acetic anhydride on aurin. A white compound easily decomposed by hydrochloric acid was obtained with the formula $\text{C}_{19}\text{H}_{14}\text{O}_3 + \text{C}_4\text{H}_6\text{O}_3$, melting at 168°. Aurin forms unstable compounds with bases, but combines with acids to form stable salts, which crystallise well. The bodies formed with acetic, hydrochloric, sulphuric, and nitric acids were prepared and examined. Aurin also combines with sulphur dioxide to form a compound,—



Rosolic acid resembles aurin in the fact that it combines with acids to form salts; the authors therefore propose to call it rosaurin. In conclusion the authors thank Mr. O'Shea, who has performed the analytical work in the paper. Some beautiful specimens were exhibited in connection with this paper.

The next paper was "*On the Derivatives of Di-isobutyl*," by W. CARLETON WILLIAMS. Di-isobutyl prepared by the action of sodium on isobutyl-bromide boils at 108°, 745 m.m., and does not solidify at -17°. Sp. gr., at 0°, 0.7088; ref. ind., at 16°, 1.3901 for the red K line refraction equivalent, 63.78. The isoprimary alcohol from the

hydrocarbon has an orange-like odour and hot burning taste. It is liquid at -17° ; boils, 179° to 180° , 765 m.m.; sp. gr., at 0° , 0.841. On oxidation it yields iso-octylic acid, an oily liquid boiling at 218° to 220° ; soluble in alcohol and ether; almost insoluble in water; sp. gr., at 0° , 0.926. The most characteristic salts are those of strontium, calcium, and copper. Ethyl-iso-octylate is lighter than water; boils at 175° . The isosecondary alcohol is obtained in smaller quantities than the iso-primary; boils at 160° to 163° ; sp. gr., at 15° , 0.820. On oxidation it yields a keton, boiling at 159° to 161° ; sp. gr., at 14° , 0.865. The keton on oxidation splits into acetic, and probably isobutyric acids.

The next paper was "On the Action of Chlorine upon Iodine," by J. B. HANWAY. The author has re-examined the question as to the existence of the compound ICl_4 , and has come to the conclusion that such a body has no existence for two reasons. First, that the reaction for its formation is impossible, as no high chloride of iodine can exist in the presence of free iodine; and, secondly, that careful experiments by which chlorine is added to iodine in the most advantageous manner for the formation of a high chloride fail to indicate such a body.

The Society then adjourned to February 6, when the discussion on Dr. Tidy's paper "The Processes for Determining the Organic Purity of Potable Waters" will take place.

CORRESPONDENCE.

A CORRECTION.

To the Editor of the Chemical News.

SIR,—I am much obliged to Mr. S. E. Phillips for pointing out, in last week's CHEMICAL NEWS, an error of which I am guilty (in vol. xxxviii., p. 259) in inadvertently putting violuric (and diluturic) acids under the head of tartaric instead of malonic acid, to which latter they seem to be more directly related.—I am, &c.,

P. T. MAIN.

January 20, 1879.

THE NEW ARSENIC TEST.

To the Editor of the Chemical News.

SIR,—If your correspondents Messrs. Otis Johnson and William Johnstone will take the trouble to examine the CHEMICAL NEWS of April 18, 1873, they will there see described by myself the reaction of aluminium and an alkali on arsenical and antimonial solutions, together with the limits of the test.

The reaction involved is certainly not new, and was first published by me as a convenient modification of the Fleitman test, Al being used instead of Zn. My pupils have constantly used the test since the above date, and prefer it in rapidity and certainty to all others.—I am, &c.,

J. W. GATEHOUSE, F.I.C.,

Public Analyst for the City of Bath.

The City Analytical Laboratory,
34, Broad Street, Bath, Jan. 17, 1879.

OXIDATIONS AND REDUCTIONS.

To the Editor of the Chemical News.

SIR,—In my table of "Oxidations and Reductions," published in the CHEMICAL NEWS, vol. xxxviii., p. 228, is an error which I desire to correct. Iodine is stated to react with oxalic acid, forming hydriodic acid and carbonic anhydride, while, in fact, no reaction occurs.—I am, &c.,

HENRY B. PARSONS,
U.S. Dept. of Agriculture.

Washington, D.C., Jan 7, 1879.

INFLUENCE OF CHLOROFORM ON NITRIFICATION.

To the Editor of the Chemical News.

SIR,—I am at a loss to understand Mr. Hehner's notion of nitrification. I have always imagined that "nitrification" implied the production of nitric acid. This production of nitric acid, according to Schloësing, Müntz, and myself is prevented by the presence of chloroform. Mr. Hehner says that nitrification is not prevented by the presence of small quantities of chloroform, and he comes to this conclusion from experiments showing that ammonia increases in amount when a small quantity of chloroform is present. This increase in the ammonia he supposes to be due to a reduction of the nitrates in the solution by bacteria. How can these experiments prove that chloroform does not prevent nitrification? Does Mr. Hehner understand by "nitrification" the destruction of nitrates, or does he suppose that the destruction and production of nitrates are allied phenomena, both of which are due to the action of bacteria?

I can only repeat, in conclusion, that chloroform prevents the formation of nitrates. No statement has been made by myself, nor, as far as I am aware, by the French experimenters, regarding its influence upon the destruction of nitrates.—I am, &c.,

R. WARINGTON

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 13, 1879.

Determination of Nitrogenous Organic Substances.

E. A. Grete.—The author finds great advantage in treating horn, wool, leather, &c., with concentrated sulphuric acid at a gentle heat previous to combustion. He suggests that the proportion of nitrogen in albuminoid bodies has been given too low.

Penta-brom-resorcin.—R. Benedikt.—On bringing penta-brom-resorcin and aniline in contact there are formed tribrom-anilin and tribrom-resorcin. Under similar circumstances phenol yields tribrom-phenol. Penta-brom-resorcin, if boiled with tin and hydrochloric acid, yields first tribrom-resorcin and then resorcin. The tribrom-reso-quinon on boiling with tin and hydrochloric acid yields tetra-brom-diresorcin.

On Mono-sulpho-lactic Acid.—C. Böttiger.—C. Schaß obtained this acid from α -chloro-propionic acid, and the author has prepared it from pyruvic acid. Both acids are identical.

On Glyoxylic Acid.—C. Böttiger.—In this fourth memoir the author describes the action of aniline upon glyoxylic acid.

Difference of the Absorption Spectra of one and the same Substance: A Reply to H. J. Moser.—H. W. Vogel.—The author points out that Moser has not given experimental proof for his assertions, and has imputed to him views which he has neither implicitly nor explicitly put forward.

Synthesis of Pyruvic Acid.—L. Claisen and J. Shadwell.—The authors prove that acetyl-formic acid, prepared from cyanacetyl, is identical with pyruvic acid.

On Dioxy-benzoic Acid.—L. Barth.—The author seeks to obtain from the formation and the behaviour of the etheroid and anhydroid derivatives of resorcin corroboration of their views on the constitution of dioxy-benzoic acid.

On Thymo-oxy-cuminic Acid.—L. Barth.—The author describes the preparation, properties, and derivatives of this acid at considerable length.

On Idrialin.—G. Goldschmidt.—The author finds this body to contain 91.71 carbon and 5.32 hydrogen, and declares that he cannot see why it should rank with the aromatic hydrocarbons.

Double Salts of Cuprous Hyposulphite (Second Memoir).—F. Kessel.—Not suitable for abstraction.

On α -Oxpara-toluylic Acid.—E. v. Gerichten and W. Rössler.—The authors pronounce this acid identical with Flesch's oxy-toluylic acid.

Action of Fuming Nitric Acid and of Nitrous Acid upon Benzol-sulphinic Acid.—W. König.—The author seeks to ascertain whether $C_{18}H_{16}N_2S_3O_6$ contains intact the three benzol-sulphuryl residues.

On Organic Thio-compounds.—O. Wallach.—The author examines the behaviour of alkyl-bromides and iodides upon sodium-thiacetamid.

New Mode of Formation of Phenyl-glyoxylic Acid.—L. Claisen and F. H. Morley.—Ethyl-glyoxylic ethyl-ether and mercury diphenyl are readily transformed with little loss into phenyl-glyoxylic ethyl-ether and mercury mono-phenyl-chloride.

On Triphenyl-methan.—E. and O. Fischer.—The authors have obtained from this compound both triphenyl-methan-cyanide and triphenyl-acetic acid. They effected this object by heating triphenyl-methan-chloride with an excess of mercury cyanide.

Action of Halogens upon the Salts of Guanidin.—J. Kamenski.—In this memoir the author examines the behaviour of guanidin carbonate with bromine and chlorine.

Chrysarobin and Alleged Chrysophanic Acid in Goa-powder.—C. Liebermann and P. Seidler.—The authors hold that the substance originally present in this powder is not chrysophanic acid, and that acid, as actually detected by Attfield, they consider a product of conversion. For the original substance they propose Attfield's synonym chrysarobin.

Synthesis of Anthrarufin and Chrysasin from Anthracen.—C. Liebermann.—On treating anthracen with sulphuric acid the authors obtained two disulpho-acids, one of which lead through an easy series of conversions to anthrarufin (Schunck and Römer) and the other to chrysasin.

Disulphanthracenic Acid and its Conversion into Anthrarufin.—C. Liebermann and K. Böck.—A detailed description of the former reaction mentioned in the previous paper.

Formulae of Rhamnetin and Xantho-rhamnin.—C. Liebermann and O. Hörmann.—The authors assign to rhamnetin the formula $C_{12}H_{16}O_5$, and to xantho-rhamnin $C_{18}H_{26}O_{22}$.

Certain Derivatives of Cœrulignon.—H. Ewald.—A description of hydrocœrulignon potassium, hexamethoxyl-diphenyl, dibrom-hexa-methoxyl-diphenyl, and dichlor-hexa-methoxyl-diphenyl.

Three Isomeric Tolidins.—A. Goldschmidt.—In addition to the tolidin obtained by Petrie the author has obtained two others. The paper contains an account of meta-azo-toluol, meta-hydrazo-toluol, and sulphate of tolidin.

On Hydrocarbons obtained by the Action of Aluminium Chloride upon Chlor-methyl and Benzol.—E. Ador and A. Rilliet.—An examination of xylole.

Determination of Phosphoric Acid as Ammonic Phospho-molybdate.—R. Finkener.—Hydrochloric and nitric acids hinder or delay the formation of the yellow precipitate, whilst dissolved molybdic acid promotes or accelerates it. Hydrochloric acid in the solution acts more powerfully than nitric, and ammoniac nitrate more powerfully than ammoniac chloride. The author's solution con-

tains per litre 33 grms. molybdic acid, 141 grms. N_2O_5 and 19.4 grms. NH_3 . When precipitating phosphoric acid the quantity of free nitric acid must always be more than enough to prevent the formation of a precipitate in the absence of phosphoric acid, but a considerable quantity of nitrate of ammonia may be dissolved in the liquid. In ordinary cases the phosphoric acid will be totally precipitated in less than twelve hours, if so much molybdic mixture is added as to make up four times the volume of the phosphoric solution, and if in every 100 c.c. of the mixture there are dissolved 25 grms. nitrate of ammonium. For washing the precipitate is employed a strong solution (20 per cent) of ammoniac nitrate, to which at first one-thirtieth of its volume of nitric acid is added. The washing is completed when the washings are no longer coloured by potassium ferrocyanide. The precipitate may be brought into a condition fit for weighing by the following operations:—After removing most of the ammoniac nitrate by means of water the contents of the filter are washed into a weighed porcelain crucible. Anything adhering to the paper is dissolved in a little warm dilute ammonia, the solution is concentrated by evaporation, nitric acid is added in excess, the whole is quickly poured into the porcelain crucible, the liquid is expelled by evaporation, and the nitrate of ammonia driven off by a flame moderated by wire gauze. The residue is hygroscopic, and must be cooled over sulphuric acid and quickly weighed in the covered crucible.

Diazo-compounds of the Fatty Series (First Memoir).—W. Zorn.—By treating ethyl-iodide with silver nitrosyl the author obtained a compound C_2H_5NO , almost as readily and as violently explosive as the chloride of nitrogen.

Discovery of Vanillin in Siamese Benzoin.—P. Jannasch and C. Rump.—This paper contains merely the analytical evidence that the substance detected was vanillin.

Preparation of Nitrous Acid.—G. Lunge.—For the preparation of nitrous acid, whether by means of arsenious acid or starch, should be used nitric acid of sp. gr. 1.30 to 1.35, and almost entirely free from hyponitrous acid.

Action of Ethyl Iodide upon Silver Maleate and Fumarate.—R. Anschütz.—The author's object is to decide the isomerism of maleic and fumaric acids by a comparative study of their derivatives.

On Alizarin Blue.—C. Græbe.—The author shows by his experiments that the correct formula for this compound is $C_{17}H_9NO_4$. By heating it with zinc-powder he obtains a new base, $C_{17}H_{11}N$, whose salts are all golden yellow, and in solution display an intense green fluorescence.

Simple Method for Obtaining the Aldehydins.—A. Ladenburg.—The author agitates dilute aqueous solutions of an ortho-diamin hydrochlorate with an aldehyd, when the corresponding aldehydin is formed with liberation of heat.

Experimental Determinations of Position.—A. Ladenburg.—Not suitable for abstraction.

On Certain Phenyl-aldehydins.—A. Ladenburg and T. Engelbrecht.—An account of phenyl-benzaldehydin and phenyl-furfuraldehydin.

On the Aldehydins.—A. Ladenburg.—The author describes dibenzyliden-amido-benzoic acid; toluofurfuraldehydin, which latter body possesses an intense bitter taste recalling that of strychnin and acts upon animals as a powerful poison; and, lastly, phenyl-anisaldehydin.

Naphtho-picric Acid and Certain of its Derivatives.—T. Diehl and V. Merz.—The authors describe the preparation of this acid and the following of its derivatives:—Amido-diimido-naphthol hydrochlorate, the chromate of the same name, and the platonic double salt.

The Formula of Uric Acid.—H. B. Hill.—The author admits an error as to the original authorship of a structural formula.

Tetra-nitroxy-sulpho-benzide.—J. Annaheim.—This compound, $C_{12}H_6N_4SO_{12}$, is a solid, straw-coloured body, of an intensely bitter taste, melting at 253° , and deflagrating at higher temperatures.

On Mucobromic Acid.—O. R. Jackson and H. B. Hill.—An account of the preparation and of certain derivatives of this acid.

Analysis of the Mineral Spring Marienbrunnen, at Huckstalle, Westphalia.—H. Vohl.

Analysis of the Bitter Water of Rákóczy, from Buda.—H. Vohl.—The interest of these two papers, if any, is medical rather than chemical.

NOTES AND QUERIES.

Gluten in Grain.—Will any of your readers inform me what grain contains the largest amount of gluten?—E.

MEETINGS FOR THE WEEK.

- SATURDAY, 25th.**—Physical, 3. "Studies in Vibration," by Dr. F. Guthrie, F.R.S.
MONDAY, 27th.—Medical, 8.30.
— Royal Geographical, 8.30.
— London Institution, 5.
— Society of Arts, 8. "Mathematical Instruments," by W. Mattieu Williams.
TUESDAY, 28th.—Civil Engineers, 8.
— Anthropological, 8.
— Royal Institution, 5. "Animal Development," Prof. Schäfer.
WEDNESDAY, 29th.—Society of Arts, 8. "The Distribution of Disease Popularly Considered," by A. Haviland, M.R.C.S.
THURSDAY, 30th.—Royal, 8.30.
— Royal Institution, 3. "Electric Induction," J. H. Gordon.
— Society of Arts, 8. "Gas Illumination," by Dr. W. Wallace, F.R.S.E.
— London Institution, 7.
FRIDAY, 31st.—Royal Institution, 9. "Logic of Architectural Design," by Mr. H. H. Statham.
— Society of Arts, 8. "Quest and Early European Settlement of India," by Dr. Birdwood, C.S.I.
SATURDAY, Feb. 1st.—Royal Institution, 3. "Reptilian Life," Prof. H. G. Seeley.

THE QUARTERLY JOURNAL OF SCIENCE.

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 - II. Gravitation as a Factor in the Organic World. By William Crookes, F.R.S.
 - III. Sanitary Science in the United States: Its Present and its Future. By Albert R. Leeds, Ph.D.
 - IV. The Course of Nature. By Prof. Simon Newcomb.
 - V. Peruvian Antiquities. By E. R. Heath, M.D.
- Notices of Scientific Works, Obituary, &c.

NOTICE.

MONTHLY ISSUE OF THE JOURNAL OF SCIENCE.

The JOURNAL OF SCIENCE will in future be issued MONTHLY instead of QUARTERLY, and will consist of 48 pp., the form and general appearance remaining the same.

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THE CHEMICAL NEWS.

VOL. XXXIX.

ON THE
3 FEB 79
ESTIMATION OF SMALL EXCESSES OF WEIGHT
BY THE BALANCE FROM THE TIME OF
VIBRATION AND THE ANGULAR DEFLECTION
OF THE BEAM.

By J. H. POYNTING, B.A., B.Sc.

WHILE working last year on an experiment to determine the mean density of the earth by the balance, I had to measure such an exceedingly small difference of weight, that I could not at that time estimate it by means of a rider, but was obliged to adopt the method described in this paper. Stated generally, it consists in treating the balance as a pendulum. Knowing the nature of the pendulum (that is its moment of inertia) and its time of vibration, we can calculate what force acting at the end of one arm of the beam will produce a given angular deflection. It is, in fact, an application to the common balance of the method which has always been used with the torsion balance when it has been necessary to calculate the forces measured in absolute measure. I cannot find any record of a previous application of the method, and as it might be of use in very delicate weighings, or in verifying the small weights in a laboratory, I have thought it worth while to give a full account of it.

When small quantities of the second order are neglected, and the oscillations are of the first order, it will easily be found that the equation of motion of the beam of the balance is—

$$(MI^2 + 2Pa^2\ddot{\theta} + (2Ph + Mgh)\dot{\theta} = a\ddot{\theta} \dots \dots (1)$$

Where MI^2 = moment of inertia of beam about central knife edge.

M = mass of beam.

a = half length of beam.

P = weight of either pan and the mass in it.

h = distance of line joining terminal knife edges below the central knife edge.

k = distance of centre of gravity of beam below central knife edge.

p = small excess in one pan.

θ = angular deflection in circular measure produced by p .

g = gravity.

If $\ddot{\theta} = \theta$ we have the position of equilibrium given by—

$$\theta = \frac{ap}{2Ph + Mgh} \dots \dots \dots (2)$$

The semi-periodic time is—

$$t = \pi \sqrt{\frac{MI^2 + 2Pa^2}{2Ph + Mgh}} \dots \dots \dots (3)$$

From equations (2) and (3) we can eliminate $2Ph + Mgh$, obtaining—

$$p = \pi^2 \frac{MI^2 + 2Pa^2}{ag} \frac{\theta}{t^2} \dots \dots \dots (4)$$

From this expression it appears that if we know the moment of inertia of the beam, its length, and the weight at each end, we can find the excess p from the time of vibration and deflection.

* Read before the Astronomer Library and Philosophical Society, December 29, 1878.

The results given in this paper were obtained with a 16-inch chemical balance by Oertling. The exact length of the half beam (a) measured by a dividing engine is 20.2484 centimetres.

To find the Moment of Inertia MI^2 of the Beam.—The simplest way theoretically would appear to be this. Find the times of vibration t_1 , t_2 and the deflections θ_1 , θ_2 due to the same excess p with two different loads P_1 , P_2 in each pan. Equating the values of p given for each by equation (4) we have—

$$\frac{MgI^2 + 2P_1a^2}{MgI^2 + 2P_2a^2} = \frac{\theta_2 t_1^2}{\theta_1 t_2^2}$$

An equation which will give MgI^2 in terms of known quantities. But on trial it was found that a very small proportional error in the observed time made a large error in the value of MgI^2 , and the following method, that usually adopted in magnetic observations, was employed in preference. A stirrup was suspended by a platinum wire, and its time of vibration (t_1) against the force of torsion (μ) of the wire was observed. The moment of inertia of the stirrup being S we have—

$$t_1^2 = \frac{\pi^2 S}{\mu}$$

The time of vibration (t_2) was then observed when a cylindrical brass bar of known moment of inertia B was inserted in the stirrup. We now have—

$$t_2^2 = \frac{\pi^2 (S + B)}{\mu}$$

The bar was then removed and the balance beam inserted in its place, and the time of vibration (t_3) gives—

$$t_3^2 = \frac{\pi^2 (S + MI^2)}{\mu}$$

From these three equations, eliminating S and μ we obtain—

$$MI^2 = \frac{B(t_3^2 - t_1^2)}{t_2^2 - t_1^2}$$

Now Bg was calculated from the weight and dimensions of the bar to be 6332.83 (in centimetres and grammes). The observed times were $t_1 = 3.6792''$; $t_2 = 4.495''$; $t_3 = 7.1483''$. From these values we find—

$$MgI^2 = 35651.6^*$$

To Measure θ .—The angle of deflection was measured by the number of divisions of the scale which the pointer moved over. As the length of the pointer is 32.1006 centimetres, while 20 divisions of the scale measure 2.5658 centimetres, a tenth of a division, in terms of which the deflection was measured, corresponds to an angle of 0.0003996. The oscillations were observed from a distance of 6 or 8 feet by a telescope. The resting-point (i.e., the point where the balance would be in equilibrium) was found in the usual way by observing the three successive extremities of two swings, and taking the mean of the second and the mean of the first and third. Five determinations of the resting-point were usually made with the excess to be measured alternately added and removed. From these five, three values of the deflection (π), due to the excess, were calculated in a manner which will be seen from the example below.

The Time of Vibration.—This was found from several determinations of the time of ten oscillations. The method will be seen from the example. No correction was needed for the resistance of the air as long as the vibrations did not exceed two divisions of the scale. When, however, they were much more than that the time of vibration was found to increase with the arc. As the time of vibration frequently changes slightly, probably through variations of temperature, it was usually observed before

* To this a small correction should be added if the adjusting bob is not in its lowest position. This amounts to 7.6 for each turn of the screw, and may therefore in general be neglected.

and after the determination of the deflection (n), and the mean of the two taken as the true time.

The following example of the determination of the value of a centigramme rider by placing it half-way along the beam will sufficiently explain the details of the method.

TIME OF VIBRATION AT COMMENCEMENT.

Pointer apparently Moving from Left to Right.

No. of Vibration.	Observed time of passage of pointer through resting-point.	No. of Vibratn.	Observed time of passage of pointer through resting-point.	Time of 10 Vibrations.
0	11h 15'36"	10	17'43"	127
2	16' 1"	12	18' 8"	127
4	16'25".5	14	18'33".5	127
6	16'52"	16	18'59"	127

Mean value of 10 vibrations 127

1	11h 15'49"	11	17'56"	127
3	16'14"	13	18'21"	127
5	16'39".5	15	18'46"	126.5
7	17' 5"	17	19'11".5	126.5

Mean value of 10 vibrations 126.75

Mean of means = 126.875.

$t = 126.875$.

DETERMINATION OF DEFLECTION n .

Excess Weight.		Extremities of Oscillation.	Resting-Point.	Mean of preceding and succeeding resting-points.	Deflection due to Excess.	
Added		109	96	102.5		
		109				
Removed..		93	40	66.25	102.25	36
		92				
Added		152	53	102	66.75	35.25
		150				
Removed..		80	55	67.25	102.5	35.25
		79				
Added		147	60	103		
		145				

Mean value of $n = 35.83$.

TIME OF VIBRATION AT END.

Pointer apparently Moving from Left to Right.

No. of Vibration.	Observed time of passing of pointer through resting-point.	No. of Vibratn.	Observed time of passage of pointer through resting-point.	Time of 10 Vibrations.
0	11h 26'19"	10	28'27"	128
2	26'44".5	12	28'53"	128.5
4	27'10"	14	29'18"	128
6	27'35".5	16	29'44"	128.5

Mean value of 10 vibrations 128.25

1	26'32".5	11	28'39"	126.5
3	26'58"	13	29' 5"	127
5	27'23".5	15	29'30".5	127
7	27'49"	17	29'56".5	127.5

Mean value of 10 vibrations 127

Mean of means = 127.625.

$t_2 = 127.625$.

Remembering that one-tenth of a division of the scale is

an angle of 0.0003996 in circular measure formula (4), expressed in milligrams., becomes—

$$p = \frac{n}{t^2} 0.3996 \frac{\mu^2}{g} (MgI^2 + 2Pa^2)$$

In our present example—

$$n = 35.83$$

$$t = \frac{t_1 + t_2}{2} = 127.25"$$

$$MgI^2 = 35651$$

$$2Pa^2 = 24704^*$$

$$p = 5.724 \text{ milligrm.}$$

The length of time occupied in this determination was not quite a quarter of an hour.

The following table contains a series of results which I have obtained of the weight of two centigramme riders, the first of which was accidentally destroyed after the conclusion of the fourth determination. As the rider was always placed at division 5 on the beam, the values given in the table are double those actually obtained:—

No. of Expt.	$MgI^2 + 2Pa^2$	t in secs.	n	Weight of rider in milligrammes.	Mean Value.
1	145364	8.921	13.458	9.78	
2	309356	17.65	25.49	10.05	
3	519769	20.435	19.12	9.55	9.96
4	130355	13.10	34.71	10.47	m.grms.
5	130355	12.87	36.6	11.44	
6	130355	12.72	35.50	11.35	
7	130355	12.725	35.83	11.45	11.35
8	130355	12.81	35.5	11.20	m.grms.
9	130355	12.903	36.37	11.31	
10	454405	19.406	22.08	10.58	

ON A NEW FORM OF APPARATUS FOR THE PRODUCTION OF FERROUS SALTS FOR TITRATION, &c.

By W. F. K. STOCK, F.C.S., F.I.C.

THE apparatus usually employed to procure the absence of atmospheric air during the solution of metallic iron in acids consists of a flask fitted with a cork, through which is passed a tube bent twice at right angles. When in use the free end of this tube is made to dip below the surface of water contained in a second flask or small beaker. The chief defects of this arrangement are:—

1st. Want of portability.

2nd. Necessity for some form of support for tube.

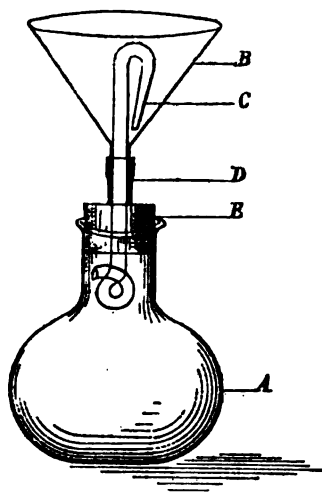
Having recently had occasion to execute a considerable number of determinations of metallic iron in samples of steel and iron, I constructed an apparatus entirely free from these defects, and very much more simple and convenient to manipulate. The following is a description of the new arrangement, which will be readily understood by the aid of the accompanying cut.

A is a flask of 200 c.c. capacity; B, a glass funnel 7 m.m. diameter; C, a glass tube of such a size as just to pass through stem of funnel B. D, an india-rubber joint connecting B and C and forming a perfectly tight joint; E, an india-rubber stopper, bored so as to pass stiffly upon C.

Perhaps a word upon the construction and use of the apparatus may be admissible. Having selected a tube of the proper size, the upper bend is turned on so that the point of the tube E, which should be narrowed to 1.5 m.m., falls into the apex of B. B is then placed in position and secured by D; the stopper E is next fitted on C, and finally the lower bend of C is turned on, taking care to allow so much tube as will prevent E being injured by heat.

* For this as for several other cases I removed the pans, and hung the weights directly by fine wires from the suspended pieces. By this means the resistance of the air was very much diminished.

In working the apparatus it is only necessary to place sufficient water in *s* to cover end of *c*, in order to secure exclusion of air. The most certain source of heat is a sand-bath. Using this I have never had a case of untimely regurgitation. Immediately the required solu-



tion is effected, the apparatus is removed from the source of heat, and recently-boiled water is run into the funnel *s* to the required extent.

Laboratory and Assay Office, Darlington,
January 22, 1879.

THE SEVEN FUNDAMENTAL TYPES OF ORGANIC CHEMISTRY,

AS VIEWED AND INTERPRETED FROM THE STANDPOINT
OF THE "TYPO-NUCLEUS" THEORY.

By OTTO RICHTER, Ph.D.

(Continued from p. 39).

In order to enable the reader to comprehend the full theoretical import and significance which attaches to this hypothesis of seven fundamental types, it becomes necessary for me to direct his attention to that most complex and elaborately finished specimen of molecular architecture in which the whole of our seven types are supposed to be collectively and completely represented, and to join me in a rapid survey of its general internal structure and organisation. Let it first of all be distinctly understood that in this our model-molecule, which relatively to the three conjugate axes of space is held to exhibit the most perfect symmetry of form and atomic arrangement, the primary constituent groups are placed side by side along the whole extent of one of these axes, to which I shall henceforth apply the term "Operative Axis," because it coincides with the particular line or direction in which chemically conflicting molecules are invariably brought to attack and to react upon each other. Let all this be taken for granted, and we may readily conceive the practicability of a mode and order of grouping, wherein our seven fundamental types have each one allotted to them a certain definite portion of space, within whose limits the primary groups are symmetrically distributed in obedience to their own special laws of molecular collocation and arrangement. It is further argued that the seven portions of space just alluded to, and which in their collective capacity may not inaptly be compared to a suite of continuous closets or chambers, are each of them destined

to serve as repositories for the exclusive accommodation of that particular class of molecules only which bear upon them the stamp of the presiding fundamental type. It is upon the basis of this, to my mind, perfectly natural and reasonable hypothesis that I shall now proceed to expound the main features of this wonderfully simple, yet all-accommodating seven-chamber system, with the view of determining not only the exact relative position of each chamber in our model molecule, but likewise what are the most salient chemical properties and functions with which its respective occupants are more or less liberally marked and endowed.

Commencing with the middle chamber, and bearing in mind that in my system the principal nucleus is invariably regarded as forming the common central point of attraction, towards which all the other component groups are, directly or indirectly, made to tend and to gravitate, it stands to reason that this identical nucleus is justly entitled to become the chief and only occupant of this middle chamber. The middle chamber, which is thus shown to claim for itself a prominently central position, is supposed to be flanked on the one side by the outer conjunct chamber, which I hold to be reserved for the exclusive accommodation of that particular class of molecules which were formerly described by me as components of the envelope, but for which the more appropriate term "outer conjunct molecules" will henceforth be substituted. Again, the middle chamber is supposed to be flanked on the other side by the inner conjunct chamber, which I hold to be reserved for the exclusive accommodation of that particular class of molecules which were formerly described by me as hydrocarbon adjuncts, but for which the more appropriate term "Inner Conjunct Molecules" will henceforth be substituted. Continuing our survey, the outer conjunct chamber is supposed to be flanked by the outer subconjunct chamber, which I hold to be reserved for the exclusive accommodation of that particular set of substituted constituents which, from their chemical character and composition, are denied free admission into the outer conjunct chamber. Again, the inner conjunct chamber is supposed to be flanked by the inner subconjunct chamber, which I hold to be reserved for the exclusive accommodation of that particular set of substituted constituents which, from their chemical character and composition, are denied free admission into the inner conjunct chamber. Completing our survey, the outer subconjunct chamber is supposed to be flanked by the outer adjunct chamber, which I hold to be reserved for the exclusive accommodation of that particular class of co-ordinately allied poly-basic and poly-acid molecules, whose parent molecule is above poly-atomic alcohol, while the inner subconjunct chamber is supposed to be flanked by the inner adjunct chamber, which I hold to be reserved for the exclusive accommodation of that particular class of subordinately allied poly-basic and poly-acid molecules, whose parent molecule is a pseudo-poly-atomic alcohol.

Having now, with the aid of our model molecule, submitted to the reader a tolerably complete general outline of the internal structure and organisation of every conceivable variety and complexity of chemical combinations, strictly so-called, I shall in the next place endeavour to explain to him the precise nature and efficacy of those three fundamental agencies or principles which are supposed to over-rule and set in motion the wonderful mechanism with which this magnificent heptarchical system of types has been so knowingly and sapiently furnished.

Commencing with the nucleus type, I proceed upon the very simple and intelligible hypothesis that all the members of this class are formed by the direct union of two chemically identical but physically dissimilar elementary molecules, E_2 , and that these physical differences, supposed to be due to certain variations in the vibratory movements of their constituent atoms, are held to constitute an efficient cause for their coalition into a molecule of double the atomic weight. The class of molecules before us will, therefore, be expressed by the general symbol $2E_2$; and

it is of importance to bear in mind that the two thus firmly welded and interlaced molecules have now as a whole become endowed with sundry very marked and tangible chemical properties and functions.

Turning from this to the second or outer conjunct type, it is conjectured that all the members of this class owe their origin and formation to the union of two not differently but similarly modified elementary molecules, E_2 . These molecules will therefore likewise be expressed by the general symbol $2E_2$; but from the peculiar conditions of their generation, which invariably require the presence and co-operation of an over-ruling nucleus, it follows that, whereas the nucleus molecules possess in a high degree the character of consistency and stability, the outer conjunct molecules are, on the contrary, entirely destitute of these qualities. Hence it becomes impossible for them to exist apart from the dominating nucleus, to which alone they are indebted for their seeming firmness and ready associability. It is clear, therefore, that, in the act of being withdrawn from under the constraining influence of that powerful centre of attraction, the liberated molecules will instantly become re-converted again into molecules of the nucleus type, from which they are understood to have originally sprung, or else into molecules of the inner conjunct type, provided always that the chemical nature of their constituent atoms is not opposed to this species of molecular metamorphosis. It is, moreover, worthy of note that, while and so long as the molecules under consideration continue to subsist in the aforesaid compulsory state of bondage and confinement, they seem to be destitute of any special chemical properties and functions, whereas these same molecules, after their conversion into the corresponding subjunct molecules, are destined to play a very prominent part in the molecular economy by co-operating with the principal nucleus towards the proper conduction and consummation of that identical process of molecular substitution which, as already pointed out on a former page, became so unduly magnified and so unwisely selected by our leading theorists as an all-sufficient ground for the erection of a solid and comprehensive chemical theory.

As regards the inner conjunct type, which falls next to be considered, I have come to the conclusion that all the members of this class require for their realisation the joint action of neither more nor less than four primary and chemically identical hydrocarbon molecules, as expressed by the general formula H_2pC_2q (p and $q = 0, 1, 2, 3, 4$, &c.), a formula which is seen to include likewise the exceptional cases, when pure hydrogen or pure carbon are made to play the part of inner conjunct molecules. The cementing agent is again believed to be the physical principle which, by causing one of these molecules to become differently modified from the remainder, determines their coalition into one single and four times heavier molecule with the general formula $4H_2pC_2q$. I may observe, in addition, that the resulting simple hydrocarbon molecules share with those of the nucleus type the character of great consistency and stability, while, in complete analogy with the members of the outer conjunct type, these same hydrocarbon molecules, from the simplest to the most complex, are capable of entering into direct chemical union with a given principal nucleus, no matter whether that nucleus happens to be presented to them from without, or whether, as is frequently the case, it has first of all been moulded out of one or other of their own component hydrogen or carbon molecules. A second very striking and remarkable feature which the inner conjunct molecules share with those of the outer conjunct chamber consists in the circumstance that, thanks to their direct chemical union with a principal nucleus, the component hydrogen and carbon molecules have now acquired the singular power and faculty of co-operating with the said nucleus towards the proper conduction and consummation of that identical process of molecular substitution to which—I repeat it with emphasis—the founder of our modern school of philo-

sophy most wrongfully insisted upon attaching such an undue weight and preponderance.

As regards the two types which follow next in the list, namely, the outer and inner subjunct types, their origin and mode of formation have already to a certain extent been traced and foreshadowed in my closing remarks on the preceding types. The presiding agency is supposed to be the principle of calority or thermal principle, and the entire process of molecular substitution, which is the outcome of the practical application of that principle to molecular statics and dynamics, will be best understood from a descriptive analysis of its three principal stages. In the first stage, the particular outer or inner conjunct molecule which is destined for conversion in the corresponding subjunct molecule is first of all made to part company with the principal nucleus, and becoming instantly subjected to the powerful influence of the ever-present physical principle, it is speedily brought to remerge under the more permanent typical form of an ordinary nucleus. In this act of dissociation the newly generated subjunct molecule is supposed to move away from the principal nucleus, without, however, passing beyond the critical point where, at a high temperature, the simultaneously formed thermal bond, as I will call it, and which is now destined to serve as the only cementing link between the principal nucleus and its similarly constituted subjunct, becomes suddenly and forcibly torn asunder. It is, of course, impossible to determine *a priori* the exact number and degree of elasticity of the various kinds of thermal bonds by means of which subjunct nuclei may remain attached to their principal nucleus. There exists, however, abundance of valuable experimental evidence to prove that in this first stage of the process certain outer subjunct nuclei, like chlorine and its congeners, remain attached to the principal nucleus by means of one thermal bond only; others, again, like oxygen and sulphur, by means of two thermal bonds; while a third class of outer subjunct nuclei, like nitrogen and phosphorus, remain attached thereto by means of three thermal bonds, that being the highest number, so far as my system is concerned, which suffices for the complete elucidation and analysis of this interesting and instructive order of chemical phenomena. There exists, moreover, abundance of valuable experimental evidence to prove that inner subjunct nuclei with hydrogen for their component element remain attached to the principal nucleus by means of one thermal bond only; while subjunct nuclei, with carbon for their component element, remain attached thereto by two thermal bonds, and not at all by means of four thermal bonds, as is still persistently taught and maintained on the other side.

(To be continued).

THE CULTIVATION OF CHEMISTRY.*

By F. W. CLARKE, B.B.,
Professor of Chemistry in the University of Cincinnati.

ONCE a year it is our pleasure and our privilege to meet together, for the interchange of views upon the questions of chemical science; for the comparison of notes in our various lines of research; for mutual help, sympathy, and improvement. It is also, I suppose, a part of our work to attract public attention to the subjects that interest us, and to do what we can to secure for chemistry a wider appreciation and greater means for development. No branch of science has done more for civilisation than ours. Old industries have been revolutionised, and new ones created; things which were once the luxuries of the few have been made the daily necessities of the many; all arts and all manufactures owe tribute to the chemist. A comparatively small number of men, a majority of them

* Address before the Permanent Sub-section of Chemistry of the American Association for the Advancement of Science, at the St. Louis Meeting, August, 1878.

teachers, working only in their intervals of leisure, established the principles which have brought about these wonderful results. If small means, widely scattered and unsystematically used, have wrought such marvels, what may we not expect from the greater opportunities which a more general comprehension of the value of our labours must eventually bring?

To the members of this sub-section, the economical achievements of chemistry are familiar as household words. As we look about us in our daily lives, we see in every direction the fruits of chemical investigations. Every scrap of metal; all paints, varnishes, fats, oils, and fertilisers; every bit of glass or porcelain; every cake of soap or box of matches, embodies some improvement which chemistry has made. Our linen is bleached and our outer garments are dyed with the products of the laboratory. Whether we burn candles, gas, or kerosene, we still have chemistry to thank for nearly all there is of cleanliness, convenience, brilliancy, purity, and cheapness in the light. In many articles of food, and in a long list of medicines; in the photograph and the galvanic battery; by the conversion of waste rubbish into articles of beauty and usefulness; in short, through a vast network of improvements and discoveries, our still infant science has established its claims to recognition as a benefactor of mankind. Would that the multitudes who have enjoyed these benefits might see their sources as clearly as we do! Then would science be fostered and encouraged, where now it struggles feebly to secure a grudging and scanty support.

A single discovery of the chemist may work peaceful revolutions in many departments of labour. Pardon me if I pause to illustrate this truism by a very familiar example. Even for us it is not a waste of time to look backward occasionally, and to consider the consequences developed from one great research or invention. Such considerations may well strengthen us in our hopes for the future of chemistry.

Nearly ninety years ago, Le Blanc discovered his famous process for converting common salt into soda. No process could be much simpler than this, and yet in its ramifications it has affected every branch of civilised society. In the first place, it widened the field of labour. Certain materials were needed at the start, namely: salt itself, charcoal, limestone, and sulphuric acid; and many workmen found employment in their production. By the new demand for sulphuric acid, this important compound was rendered cheaper, and every other chemical industry was thus directly facilitated. If the familiar saying be true that the degree of civilisation to which any country has attained may be measured by the amount of sulphuric acid it consumes, then the importance of this single item can scarcely be over-estimated. Passing from the materials employed to the process itself, we find that incidentally, as a by-product, it furnishes immense quantities of hydrochloric acid at a minimum of cost, and that here again all branches of manufacturing chemistry receive direct benefit. As a final result—if, indeed, any result can properly be called final—we find that the soda, for which the process was devised, has been enormously cheapened. In 1814, soda crystals were worth about 300 dols. per ton. By 1861, the price had fallen to 22 dols.; 5000 tons a week were produced, and 10,000 labourers found direct employment. This does not include the labour engaged in furnishing materials for the process, nor that incidentally stimulated through other industries. By the cheapening of soda, other things of more generally familiar utility were cheapened also. Chief among these we may mention glass and soap, two articles in which soda is a leading ingredient. Such a reduction in the price of soda as that just indicated could not but work wonders here. As glass and soap became cheaper, the demand for them naturally increased; and hence, through Le Blanc's invention, our houses are better lighted, cleanliness has been encouraged, and the public health, because of these steps forward, has unquestionably been improved.

In short, the results of this single invention, direct and indirect, can be traced into nearly every department of human industry. These results have been exclusively beneficial. All of us share in their advantages, and no one has been injured. They have given employment to many thousands of labourers, and that without prejudice to older occupations. And thus it is in some degree with every discovery of the chemist. Each one is like a grain of corn, small in itself, and yet a germ from which may spring the food of numberless future generations. In the course of our labours, many grains may fall by the wayside and bring but small return; still, that which sinks into fertile soil will yield a thousand fold reward to the sower.

I need build no argument upon the facts I have just given. They stand before us, not only on the pages of books, but embodied in countless manufactories scattered all over the civilised world. They render life pleasanter, easier, more comfortable. They are the sources from which future discoveries shall flow, and help to make certain the steady growth of civilisation. If the general public is not interested in chemistry, it is because we as chemists have neglected a part of our duty. We have but to speak in order to command the public ear. Our work is work of national importance, and is sure in time of national recognition. Let us ask ourselves to-day how the splendid achievements of the past may be made more fruitful, and what measures and what researches will best advance the interests of our science in the future.

It is safe to assert that in every science there is some central line of growth, to which all details are subordinated, and along which its fundamental principles find their readiest and most logical development. Here lie the germs of future generalisations; whatever promise there may be of greater exactness, either in methods or in data; and all those deeper conceptions which most intimately connect a given science with other branches of thought and knowledge. Without such a main stem, each science would be but a mass of scattered details, isolated facts, fragmentary principles, with little coherence or order. True, we may fail to recognise a real line of growth, or we may follow a false clue, and yet make discoveries of great value; but without clear ideas upon this subject the highest work is impossible. In physics, the doctrines of the conservation of energy and the correlation of forces are points on the central line. In the study of organised life, the theory of evolution indicates a main stem. How is it with chemistry?

Among chemists to-day there seem to be two schools; at least practically, if not in point of abstract theory. It is almost as if the line of growth had divided, so that we can hardly tell which is the greater, and which the lesser stem. One school, represented by a large majority of modern working chemists, seems to take an interest only in the statical side of the science; its chief aim is to discover immense numbers of new compounds, and to theorise upon their constitution. Strangely enough, these chemists have devoted nine-tenths of their energy to the compounds of a single element, carbon; scarcely regarding other substances save in so far as they unite with this or with bodies containing it. To me it sometimes seems as if, in the light of their labour, chemistry was to be defined as the science of speculating upon the possible position of theoretical atoms within imaginary molecules. Do not think, however, that I undervalue the value of the work they have done. I fully recognise its importance, although I consider it unfortunate that so much time and energy should have been spent in this one line of research, to the neglect of other, and I believe greater, fields of investigation.

The other school of chemists, a school which seems to be rapidly growing in England and to be gaining industrious votaries elsewhere, may be described as essentially dynamical in its ideas. It sees in every chemical reaction three objects of study: first, the substances which enter into the reaction; secondly, the phenomena which occur during the reaction; thirdly, the substances produced

by the reaction. The second term, the term which involves all the transformations of energy, is to them of at least equal importance with the others. In this they study the play of forces attendant upon the formation and destruction of compounds, the appearance or disappearance of heat, the velocity of the change, the effects of pressure, the alterations in volume, and so on. To them, every element and every reaction is important and interesting; they strive to see each change of composition in all its relations; to study processes as well as results; to recognise the intimate connection between chemistry and other sciences. Does it not seem as if these workers were pretty nearly on the right track? Is not their method of study the deepest and broadest? Does it not really include all there is of permanent value in the other school? To me, at least, it seems as if here the great advances are to be made, and the central line of growth discovered.

This feeling is justified, I believe by history. Much of the best progress in chemical science has been made on the physical side. In truth, chemistry and physics are but one at bottom, having their roots in the same fundamental principles. Electricity cannot be studied apart from chemical considerations, neither can light nor heat. Nor are we able to understand truly a chemical operation until we know a good deal about the physical forces which it necessarily involves. Neither science can be mastered apart from the other; for they are but two great branches from one common line of growth. The molecular theory, without which modern chemistry could hardly have existed, rests mainly upon physical foundations. Apart from the laws of Avogadro and Ampère, of Dulong and Petit, of Boyle and Mariotte, of Gay-Lussac and Charles, much which is now plain and orderly to the chemist would be chaotic and unmanageable. Doubtless, at some future time, the investigations of Mitscherlich and others into isomorphism, of Kopp into molecular volumes, of Thomsen and Berthelot into thermo-chemistry, of Gladstone into refractive indices, of Harcourt and Eason, and Boguski and Kajander into the velocity of chemical changes, will lead to generalisations of equal importance with these. The workers I have named, with many others equally worthy of naming, have already done enough to give us glimpses of great laws which we are as yet unable to see in their entirety. These glimpses forward are like promises made by Nature; sometime to be fulfilled, certainly never to be broken. Were we to take from chemistry all that has been done upon the physical side, how little true science would remain! We should have left but little law and little order; scarcely more than a vast mass of scattered facts and details, unconnected with any fundamental conceptions or any definite method of scientific research. Physics, stripped of its chemical portion, might make a somewhat better showing; but even then it would be little more than a skeleton of what it now is. Neither science can be strong without the other.

One of the main objects of science is to render prevision possible. The more thoroughly our knowledge is co-ordinated, the better are we able to predict the nearer discoveries of the future, and to see what lines of research will be most fruitful. Science is continually striving after exactness; of which this prevision is one of the results. The line, therefore, which leads most directly to definiteness and precision in any department of knowledge, is evidently the true line of growth which we should seek to find and to follow. Nature marks out pathways for us, even though we close our eyes to her indications.

To-day, notwithstanding its brilliant achievements in the past, chemistry is an inexact science. From top to bottom, from beginning to end, it shows signs of imperfection. Indeed, to most minds, this incompleteness is a great charm. There is so much to be done in chemistry, so many lines of research to follow out, so many discoveries to be made, so much glory to be won and good to be accomplished, that an active mind can hardly fail of being attracted. We may well feel, with the German

philosopher, that the power to search after truth is more precious than truth itself. Were it possible for any science to become absolutely perfect, we should but idly fold our hands and enjoy its fruits, caring little for the history behind us, or for a future which could bring us nothing more. Perhaps, in the beneficence of Nature, we might even lapse into forgetfulness, in order that our descendants could enjoy the pleasure of a re-discovery. Speaking simply in a relative sense, chemistry is an eminently imperfect science. In experimental resources it is wonderfully rich; in delicate methods of investigation no other science can surpass it; but in those principles which render foresight possible, chemistry is poor and meagre. We may guess the existence of some undiscovered compound, but until we have prepared it what can we tell of its properties? A little perhaps, a very little; and that only approximately. Of its relations to the great forces of Nature, we know in advance almost nothing. We are able only to devise for it some "structural formula" which shall last for about five years, and then be replaced by another of equally perishable quality. We are, in fact, to-day but laying the foundations of a future science, which shall be to the chemistry of the present what that is to the alchemy of the past. What the chemist has already done for humanity has been done in spite of difficulties and defects; it is but a trifle compared with that which shall be accomplished in the better time to come. Enough has been done to prove the possibility of more; to encourage the investigator in his labours; to show the public that here is something worthy of study, of help, and of applause.

There are three great objects of investigation in chemistry as seen under its present aspect; three central problems upon which all else depends. First, what laws govern the transformations of energy that occur during chemical changes? Second, how do the properties of compounds stand related to those of the elements contained in them? Third, what is the nature of chemical union? These problems must be studied largely together; each one in the light of the other two. The first and second, however, are more practical in character than the third, and involve less speculation. They need for their solution severe experimental researches, the exact determination of numerical data, and reasoning of a rigidly mathematical kind. Under the third problem we encounter the chief speculations of chemistry, hypotheses which serve to suggest and stimulate investigations with regard to the other two, while at the same time dependent upon these latter for the security of their foundations. The nature of the elements, whether one or many; the truth or falsity of the atomic theory; all questions of inter-molecular structure; the validity of certain formulæ; the value of such doctrines as that of quantivalence; these are the ideas with which this problem has to do. Each one of these theoretical questions is but a special case depending for its answer upon our views concerning the main point of all. Plainly, the first thing for chemistry to do is to make sure of its foundations; using speculation only as a line by which parts of the work may possibly be helped and guided.

The only foundations which can stand the test of time, in such a science as ours, are exact, rigorous, quantitative measurements. We already know tolerably well in a qualitative way what transformations of energy occur in chemical changes; the question now is as to their definite numerical relations. How much heat appears or vanishes in any given reaction? what is the exact electromotive force of any specified couple? what laws of quantity connect actinic energy with particular combinations or decompositions? These are the questions which science now asks, and for which in a few cases we are just beginning to find answers. The physicist approaches these questions from one side, the chemist from another; eventually the two will meet, and a general solution will be found. From our knowledge of forces on the one hand, and of substances on the other, we shall become able to compute

the dynamical relations of every possible chemical change, and perhaps even to determine in advance what reactions can and what cannot take place.

(To be continued)

PROCEEDINGS OF SOCIETIES.

PHYSICAL SOCIETY.

January 25, 1879.

Professor G. C. FOSTER, Vice-President, in the Chair.

PROF. E. RAY LANKESTER and Mr. Alex. Macdonald, B.A., were elected members.

Dr. ERCK exhibited a constant bichromate of potash battery. The ordinary bichromate battery soon loses power when in use, and in order to secure a powerful constant battery to drive a small astronomical clock, Dr. Erck devised the modified form shown. It consists of a narrow lead trough, 12 ins. long by 3 ins. wide and 1 in. deep, lined along both sides with the carbon plates. The zinc plate, 10 ins. long, is immersed in the solution to the depth of an inch midway between the two carbons. A continual circulation of the bichromate solution is kept up by allowing fresh solution to drop into the cell at one end, and the exhausted solution to drop away by a tap at the other end. As the space between the two carbons is only about half an inch wide, there is merely a thin layer of solution between the positive and negative poles. The internal resistance of the cell is therefore very low when short circuited, only about $\frac{1}{2}$ ohm. To obtain the maximum current, about 8 ozs. of solution per hour should be supplied. Dr. Erck also showed a battery formed of zinc and carbon circular plates mounted on an axle, which is rotated by wheelwork, thus mechanically stirring the bichromate solution.

Dr. GUTHRIE, F.R.S., described some the results he had obtained from experiments on the vibration of metal rods or lathes fixed in a vice at one end and free to vibrate at the other. The experiments were carried on by dusting sand on the rod and observing the nodal lines formed by it when the rod was vibrated so as to give out notes determined by a monochord. Dr. Guthrie's results showed that the two final segments at the free end are equal in length to the inner segment at the fixed end. It appears from these experiments that if a free lathe, vibrating with a node in the middle but having an even number of segments, be clamped at where there is a node we alter its conditions of vibration. When the lathe is half free the end segment breaks up into two parts together equal to the segment at the fixed end. In the case of torsional vibration of the lathe the position of the longitudinal nodal lines depended to some extent on the clamping of the lathe in the vice.

Prof. FOSTER pointed out that in a natural node the direction of the tangent is varying, whereas in an artificial node it is always horizontal.

Prof. UNWIN explained that the sand accumulated at nodes because the particles when thrown off the lathe make certain horizontal excursions, which tend to move them nearer the points of repose of the lathe.

Messrs. Elliot Brothers exhibited sundry electric commutators and resistance boxes.

EDINBURGH UNIVERSITY CHEMICAL SOCIETY.

Third Meeting, January 15, 1879.

Mr. G. CARR ROBINSON, F.R.S.E., in the Chair.

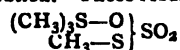
A PAPER was read by Mr. J. ADRIAN BLAIR, B.Sc., on the "Salts of Trimethyl-sulphine," containing further

results of an investigation carried on by Prof. Crum-Brown and the author.

The decomposition product of the hyposulphite of trimethyl-sulphine (CHEMICAL NEWS, vol. xxxvii., p. 130) was found by analysis to be represented by the following equation:—

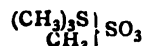


The methyl-hyposulphite of trimethyl-sulphine thus obtained is very hygroscopic, and is gradually oxidised to a sulphate. The solution of the substance does not decolourise iodine solution. These results point to—



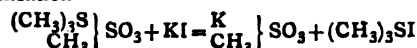
as the probable rational formula of the substance.

The sulphite of trimethyl-sulphine was obtained by the action of sulphurous acid on the hydrate. It crystallises well, but there is some difficulty in preparing a perfectly normal salt. The salt, as nearly normal as possible, does not, like the hyposulphite, give up its water of crystallisation in the cold over anhydrous phosphoric acid; at 140° C., however, it becomes anhydrous. Heated to 175° C. it gives off sulphide of methyl—8.3 grms. lost 2.32 grms., or 27.95 per cent. On cooling, the clear liquid residue solidifies, forming a hard, very hygroscopic, crystalline mass. This substance was so deliquescent that no analysis of it was made. This mode of formation leads to—

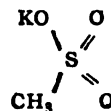


as its most probable formula.

In order to ascertain the nature of this substance the authors converted it, by double decomposition with iodide of potassium, into the corresponding potash salt, which was purified from the iodide of trimethyl-sulphine by crystallisation—

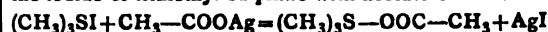


This potash salt was found to agree in properties and composition with the "sulpho-metholate" or "methyl-sulphonate" of potash.



The bearing of this fact on the constitution of mephites is obvious.

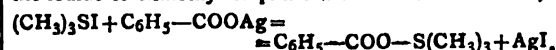
The acetate of trimethyl-sulphine is formed by treating the iodide of trimethyl-sulphine with acetate of silver.



On leaving the strong solution over sulphuric acid *in vacuo* for three weeks no crystallisation took place. The strong solution on being heated to 100° decomposed into water acetate of methyl and sulphide of methyl.



The benzoate of trimethyl-sulphine is formed by treating the iodide of trimethyl-sulphine with benzoate of silver,



This salt is very soluble in water. On standing for two weeks over sulphuric acid *in vacuo* only a very few crystals were formed, which it is difficult to separate from the very thick mother-liquor. It is slightly less soluble in alcohol. The imperfectly dried salt on being heated to 110° decomposes into water, benzoate of methyl, and sulphide of methyl.

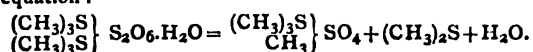


The dithionate of trimethyl-sulphine is obtained by neutralising an aqueous solution of dithionic acid with

the hydrate of trimethyl-sulphine. On evaporating the solution of the salt on the water-bath it begins after a time to crystallise out. On leaving the saturated solution to cool a large quantity of clear cubical crystals were obtained. These are not hygroscopic, insoluble in hot alcohol, and without any smell of sulphide of methyl. These properties prove it to be one of the most stable of the salts of trimethyl-sulphine. Analysis agrees with the formula—



On heating the salt to about 120° water is given off. On raising the temperature to 220° sulphurous acid is given off, and afterwards along with it sulphide of methyl, and the salt froths and melts. The heating at about 200° was continued until very little gas was given off. 8·015 grms. were found to have lost 3·325 grms., equal to 41·4 per cent. On cooling the liquid solidified. The crystalline mass was very hygroscopic, and dissolved in alcohol. On adding ether to the alcoholic solution the substance was precipitated as an aqueous syrup, and on standing over sulphuric acid crystallised out in beautiful long fine prismatic needles. The presence of the trimethyl-sulphine radical was proved by double decomposition with iodide of potassium, when the iodide of trimethyl-sulphine crystallised out. The presence of methyl-sulphuric acid was also proved. These results, along with estimations of carbon and hydrogen, prove the substance to be the methyl-sulphate of trimethyl-sulphine. The decomposition of the dithionate is therefore expressed by the following equation:—



A paper was also read by Mr. J. W. DRAKE on the "Constitution of Inorganic Salts."

NOTICES OF BOOKS.

Useful Information on Practical Electric Lighting. By KILLINGWORTH HEDGES, C.E., M.I.M.E.

A POPULAR account of the principles, production, working, and in part the cost of the electric light, confessedly somewhat hastily thrown together. Mr. Hedges divides his subject into nine sections. The introduction treats of the laws governing induced magnetic currents; the second chapter of the various kinds of electrodes and electric lamps; the third is on the division of the electric light. The kind and quantity of power required are considered, as well as the application of the light for illumination. The transmission of electricity by wires is then explained, the approximate cost of working is necessarily somewhat incompletely dealt with, trustworthy data being as yet wanting. Mr. Hedges, speaking as a practical engineer, very properly ridicules the nonsense that has been written respecting the extinction of gas by electricity. The same prophecies were made with regard to oil and candles when first gas was used. The electric light, says Mr. Hedges, will educate the public to a higher degree of artificial lighting, a prophecy already fulfilled by the Phoenix Gas Company, which has lighted up the bridge end of Waterloo Road with a brilliancy hitherto unknown in the history of gas lighting. A higher standard of gas will be called for by shopkeepers, and an increased, instead of a diminished, consumption of gas will follow. We cannot, however, agree with Mr. Hedges when he thinks that we may obtain the electric light more economically from the battery than from the dynamo-electric machine. Mr. Hedges promises to give us a second edition of his pamphlet. In the present edition he has been continually betrayed into the use of technical expressions which, although perfectly familiar to all scientific men, will be simply so much Sanscrit to the ordinary reader.

Catalogue of Chemical Apparatus and Pure Chemicals Sold by Townson and Mercer, 1879.

A WELL compiled illustrated catalogue of every description of chemical and physical apparatus. We would especially direct the attention of students, teachers, lecturers, and medical officers of health to the cheap sets of scientific apparatus described at the end of the book, which seem to be a speciality of this old-established firm. There is an excellent index at the end of the catalogue.

Practical Physics, Molecular Physics, and Sound. By FREDERICK GUTHRIE, Ph.D., F.R.S.S.L. and E., Professor of Physics in the Royal School of Mines. London: Longmans and Co., 1878.

THIS little work belongs to the excellent elementary series called the London Science Class Books now in course of publication by Messrs. Longmans. The arrangement adopted by Prof. Guthrie is somewhat new. After explaining the principles of molecular cohesion in the case of solids, liquids, and gases he carries his pupil through diffusion, effusion, osmose dialysis, and vortex motion, into wave motion in general, and lastly introduces him to the principles of sound, thus rendering the sequence complete, the string telephone and the phonograph being introduced as illustrations. There is a useful appendix added explaining the use of the vernier, the method of making parchment paper, hints on glass working, barometer making, &c. The experiments illustrating the lessons in the body of the work, and the apparatus and materials used in performing them, are also explained at greater length in a second appendix.

Notes of Statutes and Legal Decisions Affecting the Public Health Act, 1875, from 1875 to 1878. By J. V. VESSEY FITZGERALD, B.A. London: Longmans and Co.

THESE notes are intended by the author as a sequel to a former work on the same subject at the time of the passing of the Public Health Act of 1875. To officers of health, inspectors of nuisances, factory inspectors, medical men, local boards, vestries, and all other bodies or individuals whose duty it is to see that the provisions of the various health acts are duly carried out, this little work will prove invaluable.

The Retrospect of Medicine. Edited by W. BRAITHWAITE, M.D., and JAMES BRAITHWAITE, M.D. Vol. lxxviii., July to December, 1878. London: Simpkin, Marshall, and Co., 1879.

AS usual there is but little to interest the chemist in the present volume of Braithwaites' "Retrospect," valuable as it may be from a purely medical point of view. A Dr. Jencken, who hails from the Emerald Isle, recommends gun-cotton as a dressing for wounds on account of the antiseptic character of the acids used in preparing it. We should imagine that properly washed pyroxylen would have no greater antiseptic power than ordinary cleaned cotton-wool, while if it still retained sufficient traces of acid to possess antiseptic properties it would be a highly dangerous addition to hospital requisites. Of quite a different calibre is Mr. Spencer Wells's warning to surgeons—not too place to much confidence in the use of antiseptics in surgical operations at the risk of omitting any of the precautions which experience has declared to be necessary; in a word, antiseptic treatment must not be looked upon as a substitute for measures which have already proved effectual, but only as an additional safeguard. Sir Henry Thompson gives an account of the use of the microphone in sounding for stone (July, 1878), but his experiments at present do not seem to have led to any practical result. The use in psoriasis of crysophanic acid, one of the few chemical remedies discovered of late years, appears to be spreading. By the way, we may mention that a cheap method of preparing this acid

is a desideratum. It is related to anthracen, being convertible into that substance when heated with zinc-dust, the contrary operation, therefore, ought not to be impossible. The only other papers of chemical interest are those of Dr. Ogston, Jun., "On a Stale Solution of Ammonium Sulphide as a Test for Chloral; "On Butyl-chloral," by Liebreich, who finds that whereas in the case of ordinary chloral hydrate the heart is paralysed before the respiratory organs, the reverse held good in the case of butyl-chloral, a very singular and important physiological observation; and "On the Advantage of Oxygen Inhalation in Cases of Asphyxia from Breathing Carbonic Acid." Dr. R. H. Goolden announces that for thirty-five years he has used manganese sulphate with great success in the treatment of liver complaint, but that a prejudice exists against its use on account of the difficulty of getting it dispensed, pharmacutists either never having the salt in stock or taking manganese sulph. for magnesiae sulph., and dispensing it accordingly. Dr. Goolden's efforts to introduce this remedy deserve all praise, in spite of the little success that he has hitherto met with. The same remark applies to Dr. B. W. Richardson's endeavours to introduce the alkaline ethylates as caustics.

CORRESPONDENCE.

INFLUENCE OF CHLOROFORM ON NITRIFICATION.

To the Editor of the Chemical News.

SIR,—In reply to Mr. Warrington's letter, I must at once acknowledge that I used the term "nitrification" in a stretched and, I fear, erroneous sense. I certainly am of opinion, that in many cases "the destruction and production of nitrates are allied phenomena, both of which are due to the action of bacteria" or of similar organisms. Perhaps even one and the same organism induces both the formation and (under different circumstances) the reduction of nitric acid, and both these actions I included in the term "nitrification," precisely as we consider both the decomposition and the formation of carbonic acid by plants as evidences of the growth of the plant, although in reality the excretion of carbonic acid is rather a sign of decay than of increase.

When I stated that nitrification was not arrested by small quantities of chloroform I meant, and ought to have said more clearly, that chloroform is not necessarily a poison to organisms allied to, or identical, with that producing nitric acid.—I am, &c.,

OTTO HEHNER.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 27, December 30, 1878.

Reply to M. Berthelot.—M. Pasteur.

Observations Concerning M. Pasteur's Communication.—M. Trécul.—A continuation of the controversy springing out of the posthumous work of Claude Bernard.

The Microphone in Subterranean Research.—M. d'Abbadie.—The speaker, in presenting to the Academy a work by M. Michel de Rossi "Il Microfono nella Meteorologia Endogena," remarked that the author had successfully applied the microphone to the study of underground movements, and that at his observatory at Rocca

di Papa, near Rome, he had been able to distinguish three kinds of sounds, and to trace their relation with the movements of his seismometers.

Harmotome and Stilbite.—A. Gaudin.—The author gives diagrams to show the atomic arrangement of these minerals.

Electro-chemical Action under Pressure.—A. Bouvet.—The author considers that he has established the two following laws:—That the decomposition of water by a current is independent of pressure. The quantity of electricity necessary to decompose a given weight of water is the same under whatever pressure the decomposition is effected.

Liquefaction of Gases by MM. Cailletet and Pictet.—A. Bouvet.—The writer points out that the principle made use of was indicated by him in a memoir addressed to the Academy on October 8, 1877.

Magnetic Rotation of the Plane of Polarisation of Light under the Influence of the Earth.—J. Joubert.—The author admits the reclamation of H. Becquerel concerning the experiment described in his memoir of December 23.

Use of the Telephone and Microphone in Scientific Researches.—M. Hughes.—These instruments may be advantageously employed, especially in researches concerning very feeble induction-currents resulting from the movement of a magnet before a helix.

A New Electric Lamp.—E. Ducretet.—The arrangement of the lamp cannot be described intelligibly without the accompanying figure. Its principal peculiarity is the use of a mercurial column into which the points are plunged.

Existence of an Oxide of Nickel, Ni_2O_3 .—H. Baubigny.—This oxide, unlike the corresponding iron compound, is not magnetic. It is formed on treating nickel chloride at 440° with moist oxygen, and appears as a grey metallic crystalline powder. It is slowly attacked by hot hydrochloric acid with liberation of chlorine, and at a strong heat it is reduced to protoxide, losing 6.6 per cent of its weight.

Nitrates found in Beets and in some other Roots.—J. A. Barral.—The highest percentage of nitrates calculated on the dry matter is found in the roots poorest in sugar.

Inactivity of the Chrome Compounds in Comparison with Vanadium in forming Aniline-black by means of Aniline Salts in Presence of Chlorates.—G. Witz.—The author, with reference to the memoir of M. Grawitz (*Comptes Rendus*, lxxxvii., p. 844), declares that the salts of chrome cannot be used in place of the salts of vanadium in their action upon aniline salts in presence of chlorates. The presence of chrome seems even slightly injurious.

Analysis of Crude Sugars and Saccharine Matters. Determination of Water and of the Totality of Salts with Mineral Bases and Organic Acids.—E. Laugier.—The author determines moisture by desiccation in hydrogen, or in coal-gas previously purified and dried. The total mineral bases are determined by incineration in a current of oxygen at temperatures below the melting-point of chlorides. In a second portion of exactly double the weight the organic acids are extracted with ether according to Schloësing's method and half the ethereal extract is poured to the ash obtained by the former experiment, thus re-constituting the previously existing salts. The mixture is again dried in the stove as in the determination of water and weighed. The second half of the ethereal extract is then neutralised with a standard alkaline liquid. From the result obtained is calculated the weight of the carbonic acid equivalent to the organic acids. By deducting from the weight of the salts that of the ash, less the weight of the carbonic acid expelled, we obtain the weight of the total organic acids, and of the

water which their salts retain at the temperature adopted for desiccation.

Innocuity of Borax Employed in the Preservation of Food.—E. de Cyon.—The author asserts that food thus prepared preserves all its nutritive value.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 13, 1879.

Action of Dehydrating Agents upon Anhydrous Acids (Fifth Memoir).—S. Gabriel and A. Michael.—The authors examine the behaviour of phthalyl-propionic acid with sulphuric acid and with sodium amalgam; of ammonia with desoxy-benzoin-carbonic acid; and of sodium acetate with a mixture of anhydrous phthalic and isobutyric acids.

Determination of Vapour Densities.—A. W. Hofmann.—The author has been engaged with a course of experiments in order to ascertain the limits within which the determination of vapour densities in a barometric vacuum is usefully practicable. He describes a modified method for the expulsion of air, but purposes giving on a future occasion a detailed account of his methods and results.

The Formation of Methyl-aldehyd.—A. W. Hofmann.—The author passes a mixture of the vapour of methylic alcohol and of air through a platinum tube, not too narrow, containing a bundle of fine platinum wires. On the application of a gentle heat and condensing the vapours methylic aldehyd is obtained in quantity.

La Correspondance Scientifique.

December 10, 1878.

The Patent Office at Washington has refused to grant Mr. Edison a patent for his electric light, as his invention is merely a reproduction of that of Mr. Starr, of Cincinnati, whose patent dates from 1845. The English Patent Office is also said to have found Mr Edison's invention to have been anticipated by that of Chanzy, of Brussels, in 1852 laid by him before the Academy of Sciences February 25th 1858.

Syrup of orgeat is made of glucose made into an emulsion with oil and gum-tragacanth and flavoured with nitrobenzol.

The tin-paper used for covering chocolate and other essenculs has been found to contain 85 per cent of lead.

MISCELLANEOUS.

Society for the Encouragement of Arts, Manufactures, and Commerce, John Street, Adelphi, London, W.C.—One hundred and twenty-fifth session, 1878-79.—The following arrangements have been made for the forthcoming meetings of the Society:—Ordinary Meetings—Feb. 5, "The Best Methods for improving the Condition of the Blind," by Dr. T. R. Armitage. Feb. 26, "Indian Pottery at the Paris Exhibition," by George Birdwood, M.D., C.S.I. March 5, "The Social Necessity for Popular and Practical Teaching of Sanitary Science," by Joseph J. Pope, M.R.C.S., L.S.A. March 12, "The Compensation of Timekeepers," by Edward Rigg, M.A. March 19, "Economical Gardens for Londoners," by W. Mattieu Williams, F.R.A.S., F.C.S. March 26, "The Treatment of Iron to Prevent Corrosion" (a second communication), by Prof. Barff, M.A. In the Chemical Section—January 30, "Gas Illumination," by Dr. William Wallace, F.R.S.E. In the Indian Section—January 17, "Afghanistan," by C. E. D. Black, Col. H. Yule, C. B., R.E., will preside. January 21, "Quest and Early European Settlement of India," by George Birdwood, M.D., C.S.I. February 21, "The Trade of Central Asia," by Trelawney Saunders. March 7, "The Moral and Material Progress of India," by H. Phillips. In the African Section—January 21, "Retrospect and Prospect in Egypt," by B. Francis Cobb. February 4, "The Opening of the District to the North of

Lake Nyassa, with Notes of a Recent Expedition through that Country by H. B. Cotterell. March 18, "Some Remarks upon an Old Map of Africa contained in Janson's Atlas, published at Paris in 1612," communicated and exhibited by R. Ward. April 1, "The Contact of Civilisation and Barbarism in Africa, Past and Present," by Edward Hutchinson, Lay Secretary of the Church Missionary Society. **Cantor Lectures**—First Course, on "Mathematical Instruments," by W. Mattieu Williams. The remaining two lectures will be delivered on the following dates:—January 20 and January 27. The second course will be by Dr. W. H. Corfield, M.A., on "Household Sanitary Arrangements." It will consist of six lectures, to be given on the following dates:—February 17, 24, March 3, 10, 17, 24. The third course will be by W. H. Preece, on "Recent Advances in Telegraphy." It will consist of five lectures, to be given on the following dates:—April 21, 28, May 5, 12, 19. **Additional Lectures**—A Course of two lectures will be given by Dr. B. W. Richardson, M.A. L.L.D., F.R.S., on "Some Further Researches in Putrefactive Changes," in continuation and completion of his course of Cantor Lectures given last session. Any additions to or alterations in the above meetings will be duly announced.

MEETINGS FOR THE WEEK.

- MONDAY, 3rd.**—Medical, 8.30.
— London Institution, 5.
— Royal Institution, 8. General Monthly Meeting.
— Society of Arts, 8. "Some Further Researches in Putrefactive Changes," by Dr. B. W. Richardson, M.A., L.L.D., F.R.S.
TUESDAY, 4th.—Civil Engineers, 8.
— Royal Institution, 3. "Animal Development," Prof. Schäfer.
— Zoological, 8.30.
— Society of Arts, 8. "The Opening of the District to the North of Lake Nyassa, with Notes of a Recent Expedition through that Country," by H. B. Cotterell.
WEDNESDAY, 5th.—Society of Arts, 8. "The Best Method for Improving the Condition of the Blind," by Dr. T. R. Armitage.
— Geological, 8.
— Pharmaceutical, 8.
THURSDAY, 6th.—Royal, 8.30.
— Royal Institution, 3. "Electric Induction," J. H. Gordon.
— Chemical, 8. Discussion on Dr. Tidy's Paper on the Processes for Determining the Organic Purity of Potable Waters.
— Royal Society Club, 6.30.
— London Institution, 7.
FRIDAY, 7th.—Royal Institution, 9. "Bells," by Rev. H. R. Haweis.
— Geologists' Association, 8. (Anniversary.)
SATURDAY, 8th.—Royal Institution, 3. "Leasing," by Reginald W. Macan.
— Physical, 3. (Anniversary.)

THE MONTHLY JOURNAL OF SCIENCE AND ANNALS OF BIOLOGY, ASTRONOMY, GEOLOGY, INDUSTRIAL ARTS, MANUFACTURES, AND TECHNOLOGY.

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THE CHEMICAL NEWS.

VOL. XXXIX. No. 1002.

ON THE TRANSMISSION OF POWER BY MEANS OF ELECTRICITY.

By Profs. ELIHU THOMSON and EDWIN J. HOUSTON.

THE statements recently made as to the size and cost of the cable that would be needed to convey the power of Niagara Falls to a distance of several hundred miles by electricity, have induced the authors to write the present paper, in the hope that it may throw light upon this interesting subject.

As an example of some of the statements alluded to, we may cite the following, viz.: That made by a certain electrician, who asserts that the thickness of the cable required to convey the current that could be produced by the power of Niagara, would require more copper than exists in the enormous deposits in the region of Lake Superior. Another statement estimates the cost of the cable at about 60 dols. per lineal foot.

As a matter of fact, however, the thickness of the cable required to convey such power is of no particular moment. Indeed, it is possible, should it be deemed desirable, to

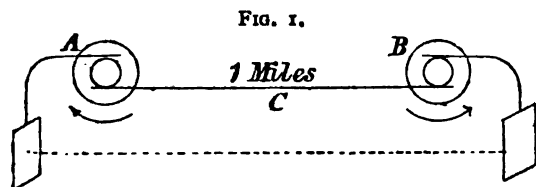


FIG. 1.

convey the total power of Niagara, a distance of 500 miles or more, by a copper cable not exceeding one-half of an inch in thickness. This, however, is an extreme case, and the exigencies of practical working would not require such restrictions as to size.

The following considerations will elucidate this matter. Suppose two machines connected by a cable of, say, 1 mile in length. One of these machines, as, for example, A,

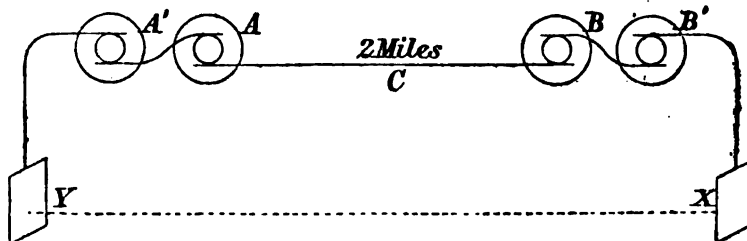


FIG. 2.

Fig. 1, is producing current by the expenditure of power; the other machine, B, used as an electrical motor, is producing power by the current transmitted to it from A by the cable C. The other terminals, x and y, are either put to earth or connected by a separate conductor.

Let us suppose that the electromotive force of the current which flows is unity. Since, by the revolution of B, a counter-electromotive force is produced to that of A, the electromotive force of the current that flows is manifestly the difference of the two. Let the resistance of A and B together be equal to unity, and that of the mile of

cable and connections between them the 0.01 of this unit. Then the current which flows will be—

$$C = \frac{E}{R} = \frac{1}{1.01}.$$

If now an additional machine, A', Fig. 2, and an additional motor, B', and an additional mile of cable, be introduced into the above circuit, the electromotive force will be doubled, and the resistances will be doubled, the current strength remaining the same as—

$$C = \frac{E}{R} = \frac{1+1}{1.01+1.01} = \frac{2}{2.02}.$$

Here it will be seen that the introduction of the two additional machines, A' B', has permitted the length of the cable c to be doubled, without increasing the strength of the current which flows, and yet allowing the expenditure of double the power at A A', and a double recovery at B B' of power, or, in other words, a double transmission of power without increase of current. Increase, now, the number of machines at A to say one thousand, and of those at B in like proportion, and the distance between them, or the length of the cable, one thousand, or in the case we have supposed, make it one thousand miles, its diameter remaining the same. Then, although the same current will flow, yet we have a thousand times the expenditure of power at one end of the cable, and a thousand-fold recovery at the other end, without increase of current. And the same would be true for any other proportion.

Since the electromotive force is increased in proportion to the increase of power transmission, the insulation of the cable and machines would require to be proportionally increased.

As an example it may be mentioned that a dynamo-electric machine used for the purpose of A in the figure, may have a resistance of say 40 ohms, and produce an electromotive force of say 400 volts. Such a machine might require from three to five horse-power when used in connection with a suitable motor B, for recovery of the power transmitted.

If the resistance of the motor B be, say, 60 ohms, and the cable transmitting the currents a distance of one mile be one ohm, then the current

$$C = \frac{400}{60+40+1} = \frac{400}{101}.$$

If, now, one thousand machines and one thousand motors, and a thousand miles of cable, each of the same relative resistances be used, the current

$$C = \frac{1000 \times 400}{1000 \times 101}.$$

which has manifestly the same value as before. If our supposition of the power used to drive one machine be correct, then from three to five thousand horse-power would be expended in driving the machines, and possibly about 50 per cent of this amount recovered. Then we have from 1500 to 2000 horse-power conveyed a distance of 1000 miles. What diameter of copper cable will be required for such transmission? Since this cable is supposed to have the resistance of one ohm to the mile, calculation would place the requisite thickness at about $\frac{1}{4}$ inch. If, however, the distance be only 500 miles, then

the resistance per mile may be doubled, or the section of the cable be decreased one-half, or its diameter will be less than the $\frac{1}{4}$ inch.

For the consumption of 1,000,000 horse-power a cable of about 3 inches in diameter would suffice under the same conditions. However, by producing a much higher electromotive force, the section of the cable could be proportionally reduced, until the theoretical estimates, which we have given in the first part of this paper, might be fulfilled. The enormous electromotive force required in the above calculation would, however, necessitate such perfect insulation of the cable, that the practical limits might soon be reached. The amount of power required to be conveyed in any one direction would, of course, be dependent upon the uses that could be found for it; and it is hardly conceivable that any one locality could advantageously use the enormous supposed power we have referred to.

Stripped of its theoretical considerations the important fact still remains, that with a cable of very limited size, an enormous quantity of power may be transferred to considerable distances. The burning of coal in the mines, and the conveyance of the power generated by the flow of rivers, may therefore be regarded as practicable, always, however, remembering that a loss of about 50 per cent will be almost unavoidable.

It may be mentioned that Dr. C. W. Siemens and Sir William Thomson have recently made statements that are in general accordance with the views here expressed.

ON THE EFFECT OF HEAT ON THE DI-IODIDE OF MERCURY, HgI_2 .*

By G. F. RODWELL, Science Master, and H. M. ELDER, a Pupil in Marlborough College.

IN continuation of the experiments on the effects of heat on the chloride, bromide, and iodide of silver, which one of us has previously had the honour of communicating to the Society,† it was thought to be advisable to search in some of the other metallic iodides for molecular anomalies similar to those presented by the iodide of silver. Among these no substance appeared more likely to possess such anomalies than the di-iodide of mercury. This substance, as is well known, is dimorphous. In the amorphous condition it presents the appearance of a brilliant scarlet powder, which, if heated, fuses at 200°C ., and volatilises just above the fusing point to a vapour more than twice as dense as that of mercury. The vapour condenses to rhombic prismatic crystals, which frequently become scarlet while cooling, but which, if they still remain yellow when cold, instantly become scarlet if rubbed or otherwise mechanically agitated. According to Warrington, this change is due to the transformation of the rhombic prisms into acute square-based octahedrons with truncated summits. If the yellow prismatic crystals are placed under the microscope, and are then touched, the change to the red variety may be observed to go on through the mass of contiguous crystals, accompanied by a slight movement, but the external form of the crystals remains unchanged, consequently pseudomorphous crystals are produced; and the larger rhombic prisms have been resolved into a mass of minute octahedrons. Frankenheim asserts that by the application of a very gentle heat both the red and the yellow crystals may be sublimed together, and he believes that the vapour of the yellow crystal passes off at a lower temperature than that of the red. Warrington found that the precipitate produced by iodide of potassium in chloride of mercury appeared under the microscope to be composed of rhombic laminæ, which gradually altered their form by the

truncation of the edges until they disappeared, while square-based octahedrons were produced in their place.

The iodide is clearly capable of existing in two crystalline forms belonging to different systems, and of passing from one form to the other, either by diminution of temperature or by simple mechanical means. Such a substance would seem to be likely to possess peculiarities in its modes of expansion under the influence of heat. In order to test this the iodide was submitted to the same experimental treatment as that employed in the case of the iodide of silver, and previously described in detail.

Homogeneous rods of the iodide of mercury were heated in paraffine in the expansion apparatus described and figured in the previous paper, and the extent of expansion due to a given range of temperature was noted. The apparatus was standardised by means of a rod of fine homogeneous silver. The same micrometer, reading to $\frac{1}{1000}$ th of an inch, was employed, and the mode of conducting the experiments was precisely the same as in the case of the iodide of silver. Two slight changes were made in the apparatus, however:—the one consisted in the substitution of a massive stone base for the wooden one hitherto used; and the other the replacement of the glass rods moving in stuffing boxes by curved equal-armed levers moving over the rim of the trough, by which means the leakage of hot paraffine at the stuffing boxes was prevented.

Bars of the iodide of mercury were cast in clean glass tubes, and here at the outset the experimental difficulties commenced. For not only was it difficult to obtain a homogeneous rod, on account of the volatilisation of the iodide at a temperature slightly exceeding its melting point, but the rod when cold was found to be so brittle that it usually broke in the attempt to remove the glass envelope from the outside. Eventually good rods were procured by slowly melting the iodide in thin glass tubes and annealing in hot paraffine. When the whole was cold the glass was cut on the outside, and carefully broken off the ends of the rod, which were sawed plane by a fine steel saw, and then furnished with metal caps, and the rod was placed between the levers of the expansion apparatus. After heating the bar once or twice in paraffine to a temperature approaching its melting-point longitudinal rifts appeared in the glass envelope, which was then easily removed, leaving a clean homogeneous rod of the iodide.

On heating a mass of the crimson amorphous iodide, it turns yellow at 126°C ., and just before the melting-point is attained the yellow changes to a deep red-brown. The liquid resulting from the fusion has the appearance of liquid iodide of silver, that is to say, it has the exact colour of bromine. The liquid when cooled solidifies to a red-brown solid, which speedily becomes yellow, and at 126°C . it changes to the crimson octahedral variety. Distinct cracking sounds, due to inter-molecular movements, were heard during the continuance of the change. Heat is absorbed when the red iodide changes to yellow, and is given out when the yellow iodide changes to the red.

A bar of the iodide was placed in the expansion apparatus, melted paraffine was forced upon it, and when the index had become quite steady, a gentle heat was applied to the paraffine. The index showed a regular and slow expansion until a temperature of 126°C . was reached, when the bar began to change from the octahedral to the prismatic condition, and without further rise of temperature rapid expansion took place. The temperature was kept constant until the change was complete, and was then slowly raised. A regular expansion now took place under a higher coefficient than before the molecular change, and this continued until the melting-point was attained. The results were concordant.

The expansion in passing from the solid to the liquid condition was determined by weighing mercury in a tube, and afterwards filling it to the same height with fused iodide. The specific gravity of each substance being

* Paper read at the Royal Society.

† *Proc. Royal Society*, vol. xxv., p. 280.

known, and the weight of equal volumes, the expansion could obviously be readily determined.

The coefficient of cubical expansion for 1° C. from 0° C. to the point of change—126° C.—was found to be:—

$$0.0000344706.$$

At 126° C., during the change from the red octahedral to the yellow prismatic condition, the body increased in bulk to the extent of:—

$$0.00720407.$$

The coefficient of cubical expansion for 1° C. from 126° C., after the change, λ , to the melting-point 200° C., was—

$$0.0001002953.$$

Thus, if we suppose a molten mass of the iodide of mercury to be cooling down from 200° C. to 0° C., the following would be the volumes under the conditions, indicated:—

Vol. at 200° C. of the liquid mass	 = 1	.1191147
" " solid mass	 = 1	.0190453
" 126° C. (yellow prismatic condition)	= 1	.0115378
" " (red octahedral condition)	= 1	.0043337
" 0° C.	 = 1	.0000000

According to Schiff the specific gravity of the octahedral iodide is 5.91; while Karsten makes it 6.2009, and Boullay 6.320.

Two distinct specimens with which we worked gave respectively—

- (1). 6.3004
- (2). 6.2941

The specific gravity of the fused iodide was found by the method before described to be

$$5.2865.$$

Thus the specific gravities corresponding to the five marked conditions shown in the curve table are as follows:—

Sp. gr. at 0° C.	 = 6	.297
" 126° C. (octahedral condition)	.. = 6	.276
" " (prismatic condition)	.. = 6	.225
" 200° C. solid	 = 6	.179
" " liquid	 = 5	.286

THE CULTIVATION OF CHEMISTRY.*

By F. W. CLARKE, S.B.,
Professor of Chemistry in the University of Cincinnati.

(Concluded from p. 51).

ALTHOUGH the foregoing statements relate specifically to the first of the three great problems, similar principles apply to the second. If we are to determine what general laws connect the properties of compounds with those of the elements contained in them, we must first secure a large mass of well-established data to work from. For a very large number of substances, such constants as density, crystalline form, melting and boiling point, thermal conductivity, specific and latent heat, co-efficient of expansion, index of refraction, electrical resistance, and so on, ought to be rigorously ascertained. With such accurate data, we may hope to reach the desired laws; without them we can see but a little way. Moreover, every constant should be determined in a much more thorough manner than any have been determined hitherto. Every measurement should be repeated many times, and under all attainable conditions; otherwise the work would be manifestly incomplete.

Fragmentary researches of this kind have already been carried out in great number; but how? In some directions

I have tried to collect the evidence, in order that we might know what data were really available, and where the worst gaps lay. The discrepancies which sometimes come to light are amazing. Take a large number of specific gravity determinations for any given compound, and see how discordant they are. The values will range through wide limits, so that it is often impossible to say which one out of several is the best. These variations may be due partly to bad work, and partly to unrecognised differences in chemical constitution. Some of them are so peculiar that they can hardly be explained, except upon the hypothesis that they spring from some undiscovered allotropy or isomerism. The metallic iodides furnish some good illustrations of this idea. The well-known properties of mercuric iodide, Rodwell's experiments upon silver iodide, and Cooke's researches into antimony iodide are sufficiently suggestive. When, therefore, we meet with discordant specific gravity determinations among other iodides, we may reasonably suppose that one experimenter has dealt with one allotrope, and another with another, both unconscious of any existing differences. But however this may be, the discordance of so many data, not only among specific gravity measurements, but in all other series of determinations, serves to show how little we really know, how much work remains to be done freshly, and how much ought to be done over again. Of all the figures upon which we now rely for the solution of these grander problems, at least three-fourths should be scrupulously re-determined; and even after that had been done our material would still be woefully insufficient. The kind of work we need may be well illustrated by the classical labours of Stas upon the atomic weights, and the immortal researches of Regnault into the physical properties of vapours. Such work as this will give imperishable foundations for exact science, strong enough and broad enough to sustain the most stately edifice which the human intellect can rear.

To my mind, the investigation of this sort most immediately needed is the accurate determination of all the physical constants for all the so-called elements. This would supply evidence directly related to all three of the great problems. We might thus learn something about the nature of the elements themselves, and how far the transformations of energy due to chemical action depended upon their individual characters; we should also be put in some position to predict the properties of compounds. In many important lines of research, science would then replace speculation, and the imagination could still find ample fields in which to exert its powers. As yet such work as this has not been done for even one of the sixty-five or sixty-six elements known. Take iron, for example,—the most important of the metals, one which has been the subject of numberless practical researches,—and we as yet know it in these more purely scientific relations barely on the surface. Not a single series of physical constants has yet been thoroughly determined with regard to it, notwithstanding the light that such investigations might shed upon important industrial problems. We know a little here and a little there; a bit about its thermal properties and another bit about its electrical properties, but nothing systematically and well. As for gold, silver, copper, and mercury, rather more is known; but even with these elements not a tenth of what ought to be done has really been accomplished.

The reason for such incompleteness in chemistry is sufficiently evident. Indeed I have already hinted at it. The labour of investigation hitherto has fallen chiefly upon the shoulders of volunteers, who have worked in great measure independently of each other, every man following his own particular bent. Part of the work has been done by teachers, in their odd moments of freedom from the drudgery of instruction; other portions were carried out by inventors, or by manufacturers with reference to special industrial questions. There has been little co-operation, and no organisation of efforts. The one class of investigators has sought for the glory which brill-

* Address before the Permanent Sub-section of Chemistry of the American Association for the Advancement of Science, at the St. Louis Meeting, August, 1878.

liant discoveries bring; the other has laboured to secure pecuniary rewards. Thus has our pure science grown unsystematically, and our applied science has been imperfectly applied. The two depend on each other; and the more complete pure science can be made, the simpler and more beneficent will its applications become.

What chemistry needs, then, is combined effort upon some general plan. There must be a body of trained workers under an efficient head, who shall mark out some of the lines for investigation to follow. Such a co-operation can hardly be brought about under existing circumstances and with present facilities, since as a matter of course every scientific man prefers to work independently, and to shape his researches for himself. Something may be done, perhaps, by those college professors who are fortunate enough to have a considerable number of advanced students to assist them; but work carried forward under such conditions can scarcely amount to more than a clearing of the ground. Still such pioneer labour is not to be despised. It marks out many useful pathways for us, and shows where the chief difficulties are to be encountered. The one great method, however, by which the desired ends may be attained is to be found in the establishment of endowed laboratories devoted to pure research. With regard to laboratories of this kind much has been written, both for and against. From two distinct points of view has the subject been treated. One set of writers has opposed the idea of endowing research, upon what appear to me to be wholly fallacious grounds. They have urged that the payment of salaries to scientific men, either as a reward for past services or as a means of encouraging future work, would probably result in the foundation of sinecures in which pretentious idlers might find an excuse for doing nothing. A man could easily pretend to make investigations, and in reality only fritter away his time in a sort of elegant leisure. No endowment would render brilliant discoveries certain, so that fruitlessness might find ready apologies. This is a somewhat extreme statement of the case, though not a wholly unjust one. Such forebodings as these might be realised, although it does not to me seem probable that the men who now labour so earnestly and assiduously for the love of knowledge would accomplish any less because of greater encouragement. Still an endowment which afforded large salaries for no regular, specific duties, might in some cases work mischief. The true man of science would hardly advise even a possible wasting of means.

The idea of an endowed laboratory for research is of a very different kind. It proposes an institution in which salaries shall be paid for work of a definite, prescribed character. Men would not be employed in it to make showy discoveries, although these might be rendered incidentally possible. The laboratory would, in short, be a place wherein the fundamental data of chemistry and physics should be accurately established, without more than casual reference to particular industrial questions or to theories. In it a body of trained specialists would co-operate upon such researches as I have suggested; they would fix some of the foundation stones of science; they would combine their strength upon those tasks which are too large and too arduous for individuals to undertake. Just as an astronomer makes an exact map of a given zone in the heavens, observing the position of each star over and over again; so the workers in this laboratory would undertake repeatedly those delicate measurements which physical science so much needs. The physical properties of substances could be exactly determined; methods of measurement could be tested and improved; apparatus too expensive for individuals to own might be constructed and employed; the quantitative relations of the several forces could be accurately ascertained. One great problem at a time might be taken up, and the energy of the whole body of workers concentrated upon it. Under ordinary existing circumstances a scientific man teaches several hours a day, and after that he labours at research. Here he would have a different routine from

that of teaching, and probably as much leisure for himself and his own ideas as before. Indeed, to carry out the latter his opportunities would be vastly improved.

Who can doubt that such a laboratory would benefit science? To have important data systematically determined, with every conceivable precaution and the best appliances, would certainly be worth while. Every industry involving applications of physics or chemistry would be aided. The special problems of the ironmaster might not be touched, and yet evidence would be discovered which should enable him to solve them much more easily and certainly than before. Precision would be gained everywhere; in facts, in methods, and in appliances; theories would rest on more solid foundations; the intellect would find new food for thought. The work done might seem at first sight to be of the hardest, driest, and most practical character; and yet there would be within it more than utility. We cannot deepen our knowledge of the real without at the same time broadening our insight into the ideal. The more we know, the vaster the unknown seems to us. The better we know, the more truly do we recognise the imperfections of our knowledge. We climb each mountain only to see a bluer, higher, grander summit far beyond.

Can this conception of a laboratory for pure research be realised? I believe it can. Physics and chemistry have created millions upon millions of wealth; they have helped to bring distant countries nearer together; they have benefited every human being within the limits of civilisation. In every country of the civilised world there are many rich men who owe their wealth to the applications of these sciences. In our own country especially, we find such men ready and often anxious to give. Every day we hear of large bequests for public purposes; of gifts to colleges, libraries, observatories, and museums; to hospitals, asylums, and churches; to the poor and to the heathen. Surely some of the wealth which our science has rendered possible ought to come back a little more directly for her benefit. There must be men who would give liberally to her purposes if they only knew of her needs. Can we not do something towards making our wants known? Are we not able to present a strong case in our own favour? Might we not by some combined effort at least make a beginning, and secure a fund, which should serve as a fulcrum for our future exertions? Let a nucleus once be established near some large university, or under the control of such an institution as the Smithsonian, and it cannot fail to grow. If the first organised laboratory, for such work as I propose, could be founded in this country, it would soon be recognised as a national glory and a benefactor of the whole human race. I throw out these suggestions merely as suggestions, with the hope that some means may speedily be found which shall lead us to the much desired end.

In Europe the needs of science are often liberally provided for by Government. The scientific institutions of Germany, for example, owe their existence and support chiefly to funds drawn from the public treasury. This policy I believe to be wise and proper, although it might not be judicious to urge it strongly here. With us, the proposal to establish a purely scientific institution, upon moneys raised by general taxation, would undoubtedly be seriously opposed. The public importance of the measure would be obscured by a clamour about economy, and upon the wickedness of expending the means of the many for the benefit of the scientific few. Much of this opposition would be foolish, although at the same time effective. Still, granting that it might be unwise to ask the National Government to establish a laboratory for research, there is another kind of laboratory which it might build, equip, and organise, with unquestionable propriety.

All the time the General Government finds it necessary to employ chemists for special expert work. For the Bureau of Engraving and Printing investigations are made with reference to the inks and paper used in the manufacture of notes and bonds. Questions are con-

tinually arising in connection with the custom houses, which can only be settled by the testimony of a reliable chemist. Frequently the Examiners of Patents would be much aided in their decisions if they had laboratory facilities at their command. The War and Navy departments are continually needing work done, it may be in the examination of supplies, in testing metals, or in experiments upon explosives. For the Lighthouse Board there are oils to be tested, and for the Geological Surveys there are rocks and minerals to be analysed. As for the Agricultural Department, it already has a small laboratory, and there is another connected with the Army Medical Museum. These examples do not by any means complete the list; they only serve to show how great an amount of scientific labour the National Government is obliged annually to employ. The work is now done in a scattered way, often with imperfect appliances, and much less thoroughly than is desirable. The question I now raise is this: Would it not be eminently proper for the Government to establish a first-class chemical and physical laboratory, in which all of its work in these departments should be done by an adequate body of scientific men, provided with the best materials and apparatus? Would not this be a measure of true public economy? Could not more and better work be done for a given sum of money than is done now? To me it seems that the plan is decidedly practical, and that Congress might by judicious pressure be induced to carry it out. Of course the measure is open to the usual political objections. It might enable inefficient men to get responsible positions through political influence; and there would be a possible danger that corruption should secure another foothold where it is already too firmly lodged. This objection I do not believe to be sound. At present experts may be employed through political favour, while the country at large knows nothing of what is going on. In a national laboratory the appointments would be conspicuous, and mismanagement would at once be recognised. There is an argument from analogy which may be even stronger than this. Whatever weakness our public officers in general may have exhibited, our scientific service has always been remarkably able and strong. If the proposed laboratory could be made as efficient as the Naval Observatory, the Coast Survey, or the Army Medical Museum; if it could secure as thorough men as these have secured, the objections just urged would disappear. The nation must get the work done somehow, and would it not be best done in the manner I propose?

Such a laboratory as this, organised at first for the merely practical work of the Government, might in course of time widen its scope a little. The National Observatory, established originally for the purposes of the Navy Department, has been able to do much for science beyond its purely routine labours. The expenditure of public money for eclipse and transit expeditions, in furnishing the means for work like that of Newcomb and Hall, has been for the interests of abstract science; but I doubt whether any intelligent citizen has ever begrudged a dollar of it. The nation has been ennobled by such researches, even though no material benefits should ever spring from them. So with a national laboratory. At first its work would be only a technical routine; but sooner or later this would necessarily develop into something more. The Government investigations could not be carried on for ever by imperfect methods; and accordingly better methods would have to be devised. Whatever researches tended towards improving the work done for the nation would, of course, be properly carried out in a national laboratory. Thus an institution, established for practical purposes at public expense, could at the same time benefit the public service, and advance the interests of science. The cost would be justified over and over again in a purely material way, so that the question of the usefulness of the laboratory could never be seriously raised.

- As we are gathered together for the Advancement of

Science, I may be permitted to broach one more subject which I believe to be of some importance. Year after year the researches of American chemists are steadily increasing in number, extent, and value. Work like that of Gibbs, Cooke, Smith, Mallet, Remsen, Jackson, and a dozen others who might be named, is certainly work of which we may reasonably feel proud. But how is all this material published? A little of it in the *American Journal of Science and Arts*; a part in certain foreign periodicals; another portion in several local transactions, as for example the *Proceedings* of the American Academy. In short, the work is widely scattered, and some of it is effectually buried beyond the reach of a majority of our fellow chemists. We need, I believe, in this country, a good chemical journal which shall fairly represent the work that America is doing for chemistry. A journal in which every chemical research published in the United States should find a place, either in full or in abstract, would certainly stimulate investigation among us, and send all over the world a truer estimate of our scientific standing. We might have for our organ a chemical periodical equal to any in the world, and I sincerely hope that we may not have to wait long for its establishment. Where or by whom it should be edited and published is a question I will not attempt to raise; suffice it to say that it ought to be controlled by no local clique, and managed in no Pharisaical spirit. It should be broad enough not to fear novelty, and courageous enough to reject trash; two qualifications more easily stated than found. I throw out these suggestions, hoping that they may take root somewhere, and that without much delay. Even though we ourselves may lack the power to carry them out, our recognition of their importance may lead to action elsewhere. If we in our gatherings can point out good pathways, others with richer means may be induced to follow them. Our united counsels and combined efforts may achieve results where, working separately, we should fail.

THE SEVEN FUNDAMENTAL TYPES OF ORGANIC CHEMISTRY,

AS VIEWED AND INTERPRETED FROM THE STANDPOINT OF THE "TYPO-NUCLEUS" THEORY.

By OTTO RICHTER, Ph.D.

(Concluded from p. 48.)

In passing on to consider the second stage of the process, we may suppose that, by a species of lateral movement on the part of the newly formed subjunct nucleus away from the outer or inner conjunct chamber towards the outer or inner subjunct chamber, that molecule, while it remains still firmly attached to the principal nucleus by means of its appropriate number of thermal bonds, will at length arrive at the particular point whence it may be brought to take up a fixed and well-defined position within the corresponding subjunct chamber, a position which it will undoubtedly continue to maintain after the whole system has returned from the nascent state of unstable equilibrium into the quiescent state of stable equilibrium. It is scarcely necessary to add that at the close of this second stage of the process no substitutional action has as yet taken place, the affected molecule having simply undergone that particular preparatory treatment which befits it for the exercise of its newly acquired chemical duties and functions. Finally, as regards the third and last stage of the process which, in the majority of cases, involves the presence and co-operation of two contending molecules, its chief characteristic feature consists in the successive interchange of one or more thermal bonds with simultaneous transfer of their respective subjuncts from one principal nucleus to the other.²

Reserving the fuller discussion of this important subject for a future communication, I shall here content myself with a brief enunciation of the general law which I hold to preside over this particular order of chemical phenomena. But before doing so it will be necessary for me to pass a few observations on the proper meaning and use of the technical terms "Multivalence" and "Quantivalence," terms which, the reader will not fail to remember, have now and then formed the subject of very animated but far from satisfactory controversies in our leading journals and debating societies.

In pondering this question from the novel and more elevated standpoint of my "Typo-nucleus" theory, I was not slow in perceiving that a fatal error had been committed by either side of combatants in not sufficiently discriminating between the absolute substitutional power and capacity of a principal nucleus pure and simple and the relative substitutional power and capacity of that same nucleus when associated with a variable number and quality of outer or inner conjunct molecules.

From an extensive series of researches in this direction I have arrived at the interesting and all-important conclusion that in the former case the said power and capacity is always measured and expressed by one single thermal bond only, because it forms the only connecting link between the two semi-nuclei into which the original molecule has been made to resolve itself, while in the latter case the said power and capacity is always measured and expressed by the variable number and quality of those outer or inner conjunct molecules which have passed from the original state of conjuncts into the typically altered state of subconjuncts. Now, in my mode of viewing the term, "Multivalence" is correctly employed whenever we wish to indicate the aggregate number of chemically similar or dissimilar conjunct molecules that have entered into direct chemical union with a given principal nucleus, while the term "Quantivalence" is correctly employed whenever we wish to indicate the number of subconjunct molecules into which the whole, or a certain part only, of associated conjunct molecules have been successively converted, it being distinctly understood that such conversion is invariably accompanied by the simultaneous formation of a corresponding number of as many thermal bonds as are lawfully required to serve as connecting links between the principal nucleus and its evolved subconjuncts.

The preceding statements and explanations will, I trust, enable the reader to grasp the full scientific bearing and significance of the remarkable law above referred to, and which may be briefly enunciated as follows:—The substitutional power and capacity of a given principal nucleus, after it has entered into direct chemical union with a given number of outer or inner conjunct molecules is a variable and fluctuating quantity, which is measured and expressed *pro tem.* by the collective number of thermal bonds emanating from that particular set of outer or inner conjunct molecules which have been made to pass from the original state of conjuncts into the typically altered state of subconjuncts."

It deserves to be borne in mind that the said power and capacity is entirely independent of the chemical nature of the element out of which the principal nucleus has been moulded, for that element is found to contribute not one single unit to the sum total of the thermal bonds collectively attached thereto. We are thus driven to the unexpected but inevitable conclusion that the substitutional power and capacity with which our modern speculators are in the habit of crediting the principal nucleus does not reside therein at all, but may be said to exist in a kind of latent state within the outer or inner conjunct molecules, until the conversion of the latter into the corresponding subconjuncts causes that power and capacity to become revealed under the form of interchangeable thermal bonds which, as a variable and fluctuating quantity, are made to range between the more or less circumscribed limits of a well-defined maximum and minimum.

It is now time for me to dismiss this singularly attrac-

tive and fascinating subject, in order to examine into the last remaining set of fundamental types which I have represented as standing under the immediate control of the principle of polarity or chemical principle. This principle rests upon the hypothesis that in their nascent or chemically active state of existence every species of molecules, from the simplest to the most complex, is capable of being transiently thrown into a state of electrochemical polarity. This new kind of agency is supposed to attain its maximum of force in the direction of a straight line intersecting the molecule at right angles to its operative axis, and coinciding with the polar axis of that molecule; while in a plane at right angles to that axis, which coincides therefore with the equatorial plane of that molecule, the said agency is supposed to attain its minimum. This hypothesis, strange and mysterious though it may appear at first sight, leads at once to the consideration of two possible alternatives. In the first alternative, any two conflicting molecules may be represented as confronting each other in such a way that the polar axis of the one becomes opposed to the polar axis of the other; while in the second alternative they may be represented as confronting each other in such a way that the polar axis of the one becomes opposed to the equatorial plane of the other.

Applying this hypothesis to the direct chemical union of two or more principal nuclei with each other, I have arrived at the conclusion that all the members of the outer adjunct type which embrace the true polyatomic alcohols and their numerous basic and acid derivatives owe their origin and formation to the first of these alternatives, and, again, that all the members of the inner adjunct type which embrace the pseudo-polyatomic alcohols and their numerous basic and acid derivatives owe their origin and formation to the second of these alternatives. It deserves special notice also that the typical differences arising from the circumstance that the chief component groups of the former class of molecules are co-ordinately, because axially and axially, united, while those of the latter class of molecules are subordinately, because axially and equatorially, united, are of such a nature as materially to interfere with the general law of molecular collocation and arrangement. From a host of well-established experimental facts and observations I have it in my power to prove that, while the general law continues to exercise an absolute control over all the members of the outer adjunct type, it is no longer applicable to the members of the inner adjunct type, where its frequent infringement is apt to give rise to a great number and variety of isomeric compounds, the formation of which seems chiefly to depend upon the precise order in which certain organic and inorganic bases or acids are made to unite with that class of molecules.

Having now submitted to the reader a brief and tolerably complete general outline of the seven fundamental types of chemistry as collectively exhibited and placed side by side in our model molecule, I may yet be allowed, in drawing to a close, to give vent to a few appropriate remarks and sentiments.

It is my humble opinion—an opinion that is probably shared by the majority of unbiassed and more calmly reflecting chemists—that, in the course of its marvellously rapid progress and development our science has at length arrived at that critical turning-point where its votaries are compelled to answer to themselves the question—"Is it meet and desirable in the face of such an overwhelming mass of ever-accumulating facts that we should still go on contenting ourselves with the adulterated and unpalatable compost of antiquated notions and untenable dogmas, hallowed though they may be by the honoured memories of so many deserving and illustrious experimentalists, and may it not be that our boasted knowledge and pretended insight into nature's secrets is merely a gross delusion and a snare, the natural result, in fact, of grave speculative errors and prejudices, which must inevitably melt away under the genial rays of a more profound and enlightened

system of chemical philosophy?" Already, if we may judge from certain highly interesting and invaluable contributions to spectroscopy with which Mr. Norman Lockyer and others have recently regaled the chemical profession, a new and unexpected light has suddenly been brought to bear upon the all-important question whether the chemical elements ought still to be regarded as simple bodies in the ordinary sense of the word, or whether these elements ought not rather to be regarded as more or less complex bodies that, under the dissociating influence of temperature, may, like other truly compound substances, be brought to resolve themselves into molecules of lower atomic weight, and presumably also of different chemical and physical properties, from their parent molecules. It is contended—and I think rightly so—that these wonderful spectroscopic discoveries are totally inexplicable on the prevailing biatomic hypothesis. What other hypothesis then, I would ask the reader in all humility, remains for our final adoption but that same polyatomic hypothesis which forms the theoretical basis of that very system of chemical philosophy I have been the first in expounding and elaborating—a system, moreover, which, with respect to the splitting up or condensation of chemically simple or compound molecules, presents no difficulties whatsoever, while it possesses in these two opposite tendencies a ready and fertile source for the complete elucidation of many hitherto obscure and unintelligible chemical phenomena. With regard to the fundamental axiom which underlies my whole chemical system, namely, that not only every species of chemically elementary molecules, but likewise every variety of chemically compound molecules, as embraced under the generic term "Model Molecule," are made to exhibit relatively to the three conjugate axes of space the most perfect symmetry of form and atomic arrangement, I have yet to observe that our model molecule with its seven-chamber system, in order to satisfy the aforesaid fundamental law, requires to be doubled; in other words, that such molecules require to be united into one by joining them together with their inner adjunct chambers. It will at once be seen that, under this new aspect, the symmetry of the whole system is rendered perfect; and I may now state, in addition, that this re-duplication of our model molecule will not oblige me to alter my customary mode of notation, if it is taken for granted that any molecular changes in the right moiety of our model molecule are invariably and simultaneously accompanied by similar molecular changes in the left moiety.

In now taking final leave of my subject, I may yet be permitted to express an anxious hope that the arduous but imperatively necessary task of a thorough revision and reconstruction of the groundwork of theoretical chemistry—a task initiated by me a number of years ago, and evidently desired and prayed for by that small but rapidly increasing band of chemical practitioners whose reasoning powers have not as yet been crippled and paralysed under the withering influence of a perverse and unyielding dogmatism—may soon engage the earnest and undivided attention of our leading men of science, and induce them to countenance and second my isolated and hitherto scarcely appreciated efforts by honouring with a fair, impartial, and candid criticism those novel and original views and principles I have ventured to advance and to publish in this and preceding communications.

A NEW FAST GREEN AND MALACHITE GREEN.

By Messrs. BINDSCHIEDLER and BUSCH.

Up to the present the so-called methyl-green has been obtained from dimethyl-aniline by converting the latter into violet by oxidation, and then the violet again by methylation into green.

It has long been the endeavour of the manufacturers of aniline colours to get the green direct from dimethyl-

aniline without going through the violet; and taking as starting-point the theoretical and meritorious studies of Prof. A. Baeyer, Otto Fischer, in Munich, and Oscar Doebner, in Berlin, it has been found possible to produce green direct from dimethyl-aniline, and even at a very low price.

These direct greens are brought into the market under the names of New Fast Green, Malachite Green, &c., and possess about the same chemical qualities, but the mode of manufacture and the attention paid to the same, cause more or less a difference in brilliancy of shade and yielding. It is, however, very easy to distinguish direct greens from methyl-greens through the simple fact that wool, silk, or cotton, when dyed with direct greens, can stand a very high degree of heat without the green colour turning into violet; whilst stuff dyed with methyl-greens turns violet when brought into contact with an iron exceeding 120° of heat.

The aforesaid theoretical mode of Otto Fischer, in Munich, was published in the reports of the Berlin Chemical Society on August 11, 1877, under the title of "Condensations Produkte Tertiärer Aromatischer Basen." Otto Fischer describes therein a new base, which he obtains by the reaction of dimethyl-aniline, chloride of zinc, and hydride of benzyl, and further states that oxidising the salts derived from his method led to the production of a beautiful bluish green colour.

The exceedingly high price of the hydride of benzyl has hitherto prevented the practical application of Fischer's most interesting discovery. Now, however, this material is produced at a price which renders its practical use quite possible.

The process of manufacturing malachite-green is based upon the theoretical studies of Oscar Doebner, published in the reports of the Berlin Chemical Society, May 27, 1878.

Doebner proves in his report that green can be obtained direct from dimethyl-aniline, chloride of zinc, and trichloride of benzol.

Besides the before-mentioned system of producing direct greens through the medium of hydride of benzyl, there are other methods by which the same can be obtained: for instance, the product of the reaction of dimethyl-aniline, chloride of zinc, and dichloride of benzoyl, submitted to oxidation, gives a very beautiful and cheap green.

From the preceding statement it is obvious that different methods of manufacturing direct green exist.

THE GRAHAM LECTURE.

THE Philosophical Society of Glasgow have determined to institute a triennial lecture in memory of the late Prof. Graham, Master of the Mint, who was for thirty or forty years a member of that Society. He communicated some of his earliest papers to it before publication of its Transactions was undertaken. The Council have also caused a portrait medal of the value of £10 to be struck, and this they propose to award every third year for the best original investigation, either in chemical physics or in pure or applied chemistry, which may be communicated to the Society or to its Chemical Section in the interval of the awards. The choice as first Graham Lecturer appropriately fell on Mr. W. Chandler Roberts, F.R.S., who had so long been associated with Graham's work, and who was enabled to employ for the purposes of demonstration much of the apparatus that had been used by his distinguished master. A very large audience assembled on the 22nd of January in the Hall of Glasgow University, where Graham graduated in 1824, and the chair was at first occupied by Mr. JAMES MACTEAR, President of the Chemical Section, who pointed out that they were doubly fortunate in having secured the services of Mr. Roberts, and in the fact that Mr. James Young, F.R.S., of Kelly,

the life-long friend of Graham, had consented to preside on the occasion; he therefore vacated the chair in favour of Mr. YOUNG, who briefly introduced the Lecturer.

Mr. ROBERTS briefly traced the influence of Black and Thomson in turning the attention of Graham to the study of molecular physics, to which he patiently devoted his life. In connection with the law of diffusion of gases, the Lecturer claimed that Priestley made in 1799 an observation on the escape of hydrogen from a cracked jar. The subsequent and independent discovery of this phenomenon by Doeberiner in 1823 has hitherto been considered the starting-point of the experimental study of gaseous diffusion, to which it undoubtedly attracted Graham's attention. After a brief notice of the influence of Eastern and Greek thought on the study of molecular movement, allusion was made to Sir Christopher Wren's model representing the effects of all sorts of impulses that result from the impact of hard globulous bodies, which, according to Dr. Sprat (historian of the Royal Society), he proposed as the principles of all demonstrations in natural philosophy, it being considered "that generation, corruption, and all the vicissitudes of nature are nothing else but the effects arising from the meeting of little bodies, of different figures, magnitudes, and velocities." Herapath's revival of Bernoulli's view as to the movement of gaseous particles was considered, and Mr. Roberts then described in detail the experiments that enabled Graham to establish the law of the diffusion of gases, and he illustrated experimentally the passage of gases through porous bodies, such as unglazed earthenware and artificial graphite, as well as through a layer of the hard translucent variety of opal known as hydropbane. The mode in which Graham studied the diffusion of the momentum of gases, by observations on viscosity as indicated by rates of flow through capillary tubes, was then described. It was pointed out that his law of diffusion forms the basis of the science of molecular mechanics, and his measurements of the rates of diffusion prove to be the measure of molecular velocities which have been so profoundly investigated mathematically by Clerk-Maxwell, Clausius, and Boltzmann, and experimentally by Loschmidt in developing the dynamical theory of gases. Mr. Roberts then considered the passage of gases through colloid or jelly-like bodies which have no sensible pores, dwelling more especially on the separation of oxygen from air by the transmission of air through a thin film of india-rubber, a circumstance of special interest from a physiological point of view. The liquefaction of gases formed the subject of one of Graham's earliest papers in 1826, and it occupied his attention at intervals during his life. He held the view that hydrogen when absorbed by palladium is reduced to the metallic form, a supposition which has received strong confirmation from the success that has attended M. Raoul Pictet's efforts to solidify this gas; and that distinguished physicist stated in a letter to Mr. Roberts that it is probable Graham's indication of the density of solid hydrogen will prove to be nearly correct. Allusion was then made to Graham's opinion that the various kinds of matter now recognised as different elementary substances may possess one and the same ultimate or atomic molecule existing in different conditions of movement, the varying degrees of rapidity of this movement constituting, in fact, the difference between the elementary bodies. In other words, if the molecular energy of a so-called element could be changed, the element would be dissociated, a view of special interest in relation to the researches of Lockyer. The lecture was illustrated by many effective experiments, and concluded with the statement that it had not been instituted from the merely special interest of Graham's researches to the physicist and chemist, but in honour of the labours of a life, the memory of which will be as enduring as its work, and to stimulate others to investigate as patiently and earnestly the varied phenomena whose basis is "Molecular Mobility."

— Sir WILLIAM THOMSON, in proposing a vote of thanks

to the Lecturer, called attention to a diagram on the wall recording the rates of passage of gases by diffusion, effusion, transpiration, and by the peculiar passage through such "colloid septa" as non-crystalline metals or india-rubber; and he stated that before Graham's time these valuable physical constants were absolutely unknown. They had listened with much interest to the connection which had been traced between Graham's law of diffusion and the science of molecular physics, as well as to the account of Graham's work generally, so carefully set before them by Graham's pupil and friend.

PROCEEDINGS OF SOCIETIES.

MANCHESTER LITERARY AND PHILOSOPHICAL SOCIETY.

Ordinary Meeting, December 24, 1878.

J. P. JOULE, D.C.L., LL.D., F.R.S., &c., President, in the Chair.

'Note on the Intensity of Moonlight,' by HARRY GRIMSHAW, F.C.S.

At the meeting of this Society held on the 26th of November, 1878, some remarks were made upon the relative intensity of moonlight, daylight during a thick fog, and the diffused light of the electric arc.

The opinion was then expressed by some members that it was very doubtful whether it was possible to read printed type of anything like a small size by the brightest moonlight in this country.

The light of the moon on the 8th of December being of considerable intensity, I took the opportunity of making the experiment, which I have previously, at different times, performed more roughly, with sufficient accuracy to form some standard of comparison. The result was that I found it possible to read with certainty a still smaller type than I had previously known it possible to peruse.

The types on which I experimented were those of a newspaper, and known technically by the printers as "minion," "bourgeois," and "nonpareil." These are all smaller than the letters of the *Proceedings* of the Society, which I believe is called "small pica." The first two are what are ordinarily used for such printing as that of newspapers and the cheaper periodicals, whilst the last ("nonpareil") is a very small letter indeed. The last paragraph of this note is printed in the type in question. The printed matter in the "minion" and "bourgeois" letter was read with comparative ease; the "nonpareil" had to be perused slowly, but could be made out with certainty.

The time at which the above experiment was performed was 8.15 p.m., and the evening of course bright and cloudless. The moon was full at 7.49 p.m. the following evening. The observers were three in number, who all succeeded in reading the three types, and it should perhaps be added, as a quantity affecting the results, that the "eyes" were all comparatively young, being under the age of thirty.

CORRESPONDENCE.

WATER ANALYSIS.

To the Editor of the Chemical News.

SIR,—I have perused with much interest Dr. Tidy's valuable paper on "Water Analysis," published in the January number of the *Journal of the Chemical Society*, and am of opinion that he has done good service in drawing attention to the most accurate methods to be adopted in the examination of waters, more especially to the employment of the two-foot tube and the action of permanganate during the first hour.

The point which still remains to be discovered, however, is where to draw the line; this is the main question, and

is by no means decided by any paper on water analysis yet published. For years past our text-books, both English and foreign, have stated that when the total impurities of a sample of water reach beyond a certain figure the water is to be condemned. But this is a mere assertion which has been copied from one book into another for the last thirty years or more. The fact is, we possess at present no positive data whereby to condemn a sample of drinking-water, and cannot possess such data until physiological experiments have been made to prove when and why a given kind of water is bad or good. In most cases it is not difficult to prove a water good for drinking, but quite the reverse to establish that it is injurious to health or unwholesome, unless it be very bad indeed.

What the chemist requires is to be able to state with certainty that a water which discolours so much permanganate in an hour is unwholesome. How much permanganate of a given strength may a water discolour and yet be good for drinking? That is a point of the utmost importance, which Dr. Tidy's laborious and interesting investigations have not solved any more than his predecessors' in the same branch of research have done. Of the hundreds of analyses which pass through my hands every year there are none which give me so much trouble and anxiety (I can use no other word) than drinking-waters. Let us suppose that the ingredients of a well-water consist of carbonate of lime, 25 grains; sulphate of lime, 18 grs.; nitrates and chlorides of magnesium, calcium, and sodium, 20 grs.; organic matter, 7 grs.; total, 70 grs.; and that the microscope shows neither microphytes nor animalculi in the water or slight deposit it may leave on standing; that the 7 grains of organic matter consist almost entirely of crenate of ammonia which can be separated by alcohol. Is such a water unwholesome to drink? If so, why? Now let us suppose that these 7 grains of organic matter are some putrescent matter in a state of decomposition. Would the water then be injurious to health? I must confess that we have no positive answer to this important question. (And 7 grains of organic matter is more than is present in the great majority of waters used for drinking.)

The thing to be determined is, as I said before, where to draw the line. On what authority are we to condemn a water as unfit for drinking? Perhaps the permanganate solution as used by Dr. Tidy may help to solve the problem. The questions which require to be solved are apparently very simple:—

- (1.) How much organic matter in a putrescent state (*i.e.*, how much permanganate in the hour) may be tolerated in a specimen of water before that water is condemned as unfit for drinking?
- (2.) How much nitrates and chlorides may be present in a gallon of water before this water is pronounced unwholesome?

These questions cannot be answered by mere supposition or conjecture.

Let me recommend these points to medical men who are devoting themselves to the examination of drinking-water.—I am, &c.,

T. L. PHIPSON, Ph.D., F.C.S., &c.
Laboratory of Analytical Chemistry,
Putney, London, S.W.
February, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 1, January 6, 1879.

Reply to M. Pasteur.—M. Berthelot.—The author points out that the posthumous papers of Claude Bernard

contain merely the beginnings of a series of experiments. The question was not to open a polemic on a train of research interrupted by the death of its author, but to preserve the trace of it in science. As to M. Pasteur's theory that certain organisms derive combined oxygen from sugar instead of free oxygen from the atmosphere the burden of proof rests with him and not with dissidents.

Formation of Organic Ultramarines.—M. de Forcrand.—The author concluded that it might be possible to obtain organic ultramarines by heating silver ultramarine with the chlorides or iodides of the alcoholic radicles. He caused an excess of hydriodic ether to react upon silver ultramarine for fifty to sixty hours at 180° in a closed vessel, and obtained thus an ethyl-ultramarine in the form of a grey powder.

Separation of Ethyl-amines.—E. Duvillier and A. Buisine.—By the process which the authors have adopted they succeed in separating di-ethyl-amin as a perfectly crystalline body. The methyl-amins appear to act in the same manner.

No. 2, January 13, 1879.

Researches on Ozone and the Electric Effluve.—M. Berthelot.—Hydrogen and oxygen mixed in the proportions of two vols. of the former to one of the latter, do not combine under the influence of the effluve, whether in sealed concentric glass tubes or in a tube surrounded with a lamellar spiral of platinum and placed over mercury. The tension of these experiments was about equal to that which gives sparks of 7 to 8 centimetres in length through air when operating with an induction coil fitted with condensers. Doubtless on increasing the tensions progressively nearly up to those which produce disruptive discharges water would be formed. But it is important to show that this formation does not ensue with tensions such as the above and in conditions where a very notable quantity of ozone is produced. The resistance of hydrogen to combination under these circumstances is so much the more remarkable as the conditions are precisely those in which oxygen combines with metals, with sulphurous and arsenious acids, iodine, and even with nitrogen. Vapour of water is not decomposed by the effluve in these conditions, and oxygen does not combine with water to form hydrogen peroxide. On the other hand, carbonic oxide and oxygen combine under the influence of similar electric tensions, though even in presence of an excess of oxygen a portion of carbonic oxide remains uncombined. Reciprocally it does not prevent the incipient decomposition of carbonic acid, and in a mixture of equal volumes of carbonic acid and oxygen after twelve hours 5 per cent of the gas was decomposed into carbonic oxide and oxygen. This oxygen contained a very strong proportion of ozone or of percarbonic acid. The decomposition of pure carbonic acid by the effluve, effected in a space free from mercury and from oxidisable bodies, gives rise to special phenomena very worthy of interest, as they lead us to suspect the existence of percarbonic acid. The gas formed attacked mercury with extreme violence. It may be considered either as oxygen very rich in ozone or as containing a strong dose of percarbonic acid, C₂O₆.

Formation of Ethers of Hydracids in the Gaseous State.—M. Berthelot.—The author gives some new determinations to permit the calculation of the heat disengaged by the gaseous products of the union of the carbides of hydrogen with the halogens and the hydracids.

Are there among the Lower Species with which we are Occupied Species Exclusively Aerobian and others Exclusively Anaerobian? Are all these Beings to be Ranked in Two Classes, or in Three as M. Pasteur has Successively Admitted, or in One, as I have Recently Indicated?—M. Trecul.—The author concludes that organised ferments are merely particular conditions of species more or less complicated, which are modified according to the media they inhabit.

Instead of three classes of inferior beings, according to the present views of M. Pasteur, we must recognise one only, each species being able to present one or more ærobian and one or more anerobian conditions.

Second Reply to M. Berthelot.—M. Pasteur.—A continuation of the controversy springing from the posthumous writings of Claude Bernard.

Researches on the Compressibility of Gases.—L. Cailliet.—The author describes the improved manometer used in his recent researches. He finds that nitrogen is compressed at first more than is indicated by Mariotte's law, and that its compressibility then decreases, as he found to be the case with atmospheric air in his earlier researches. Nitrogen presents this curious maximum at about the pressure of 70 metres of mercury.

A New Compound Prism for Direct Vision Spectroscopes, having a very great Dispersive Power.—A. Thollon.—The sulphide of carbon prism presented to the Academy possesses an extraordinary dispersive power. Instead of being enclosed laterally by plates with parallel surfaces, it is enclosed by prisms of crown glass whose refrangent angles are in an opposite direction to that of the sulphide. The refrangent media are distributed as in the prism of Amici, with this difference, that the angles of the crowns are smaller, and that at entrance as well as at exit the luminous ray passes always between the summit of the angle and the normal at the surface. This system is intermediate between the simple prism and the direct vision prism, the deviation being less than in the former and the dispersion greater than in the latter. The absorption is almost annulled.

Thollon's Spectroscope.—L. Laurent.—A further description of this apparatus. Its dispersion is equivalent to that of four flint prisms of mean density.

Synthesis of Uric Derivatives of the Alloxanic Series.—E. Grimaux.—The direct product of the action of phosphorus oxychloride upon a mixture of malonic acid and urea is malonyl-urea, the barbituric acid of Baeyer, mixed with a yellow amorphous substance, sparingly soluble in water.

Action of Diastase, Saliva, and Pancreatic Juice upon Starch and Glycogen.—F. Musculus and J. de Méring.—Saliva and pancreatic juice yield with starch the same decomposition products as diastase, namely, reductive dextrines, maltose, and glucose. Glycogen yields similar products, but its dextrines are less hygroscopic than those of starch, and their reductive power is less. There is only one glycogen, whether the animal producing it has been fed exclusively on the hydrates of carbon or on albuminoid food. The existence of reductive dextrines of variable power accompanying maltose and glucose proves the necessity of having recourse to fermentation in the determination of sugar in industrial products, and explains the discordant results obtained by experimentalists. [This memoir is, curiously enough, placed under the heading "Inorganic Chemistry."]

Les Mondes, Revue Hebdomadaire des Sciences.
No. 15, December 12, 1878.

M. Maurice Perrin points out that the physiological effects of chloroform at present are much less satisfactory than was the case some years ago. He considers that less care is employed in its manufacture.

The process adopted by M. Coquerel, of the Clichy Manure Works, is said to have succeeded in instantly converting excrementitious matters into an inodorous cake, containing 3 per cent of nitrogen and 10 per cent of phosphoric acid.

Note on the Phylloxera.—M. Carves.—The author maintains that the Phylloxera *per se* is not the cause of the death of the vine, the real enemy being a fungus which plants itself in the wound made by the animal.

MISCELLANEOUS.

The Royal Institution of Great Britain.—At the General Monthly Meeting held on Monday, February 3, 1879, Warren De la Rue, D.C.L., F.R.S., was elected Secretary of the Royal Institution, and William Spottiswoode, D.C.L., Pres. R.S., was elected Manager.

NOTES AND QUERIES.

Ammonia Tables.—Can you inform us if there are any tables published of liquor ammonia strengths, taking into consideration the different temperatures as well as the specific gravity? For instance, if liq. am. is 0.895 at 35° F. what sp. gr. will it be at 60° F., and *vice versa*? We find that all the tables in Watts, Ure, and Miller always suppose the temperature to be 60°, but as we bottle at all temperatures it would assist us if we had a table that could be relied upon.—W.R.E.

MEETINGS FOR THE WEEK.

SATURDAY, 8th.—Physical, 3. "Variation of the Thermal Conductivity of a Rod with Temperature," Dr. O. J. Lodge. Annual General Meeting.
MONDAY, 10th.—Medical, 8.30.
London Institution, 5.
Royal Geographical, 8.30.
TUESDAY, 11th.—Civil Engineers, 8.
Photographic, 8. (Anniversary).
Royal Institution, 3. "Animal Development," Prof. Schäfer.
Anthropological, 8.
WEDNESDAY, 12th.—Society of Arts, 8.
Microscopical, 8. (Anniversary).
THURSDAY, 13th.—Royal, 8.30.
Royal Institution, 3. "Sound," Prof. Tyndall.
Royal Society Club, 6.30.
FRIDAY, 14th.—Royal Institution, 9. "November Meteors," Prof. Johnstone Stoney.
Quekett, 8.
SATURDAY, 15th.—Royal Institution, 3. "Lessing," by Reginald W. Macan.

TO CORRESPONDENTS.

J. A. Fahie.—We regret we cannot give our correspondent the information he requires.
S.—You had better write direct to the Professor and ask the question.

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THE CHEMICAL NEWS.

Vol. XXXIX. No. 1893.

THE DISSOCIATION OF THE ELEMENTS.

It has fallen to the lot of modern chemists to witness a discovery in their science which, if duly verified, must be pronounced of importance almost beyond parallel. The formation of endless compounds, the speculations on their "constitution," the synthesis of organic bodies, the detection of new metals,—all, in short, that can be done with or deduced from the isolation and combination, the arrangement and re-arrangement of the so-called elements, falls into a secondary position in comparison with the complete reversal of our accepted notions concerning those elements themselves. It is therefore incumbent upon us all to rise to the level of the occasion. We must neither, in view of the dazzling prospect thrown open before us, give too free rein to the "scientific imagination," nor should we carry scepticism further than candour sanctions and than the interests of Science require.

It can scarcely be denied that some among us have sinned in both these directions.

In considering the evidence laid before us in support of the decomposition of our present elements, we must remember that the discovery does not break upon us unannounced and unprepared. Whoever has heedfully dwelt on the philosophy of chemistry must admit that, as research has been pushed further and further, the primary and indecomposable character of those sixty odd bodies which we have been taught to call simple has been rendered more and more doubtful. If we consider their comparative properties we find groups which present such features as might be expected from a compound formed by some body, x , with successively increasing proportions of another, y . But we should be compelled almost to write the history of chemistry from the beginning of the present century were we to enumerate all the phenomena and all the considerations which caution us against ascribing to the "elements" more than a merely provisional character. As Mr. Lockyer judiciously points out, with the application of increasing temperatures we have increasing dissociation, and on the faith of the principle of continuity we may expect that still higher temperatures will act in the same direction as lower ones have already done. The only reply is, that hitherto we have had no satisfactory proof that this has been done in any one instance. This difficulty Mr. Lockyer's recent researches seem to have overcome. It is a matter of some importance to note that these researches were not undertaken in the hope of decomposing the elements. The object in view was a knowledge of the chemical nature of the sun, in pursuit of which the observer was compelled to form his present conclusions. It will not be necessary or us to recapitulate the evidence which has been given in detail in the CHEMICAL NEWS for January 3rd and 10th of the present year. We will merely cite the experimental conclusions that "the common lines visible in the spectra of different elements at high identical temperatures point to a common origin, and that the different lines visible in the spectra of the same substance at high and low temperatures prove that at the former dissociation goes on as continuously as at all lower temperatures."

All this, however, it may be objected, is not direct experimental evidence, analytic and synthetic, such as chemists are able to produce in favour of the compound nature of water. Mr. Lockyer has not, *e.g.*, taken a gramme of copper and decomposed it, exhibiting, in a state of isolation and in due proportion, each of the various more elementary elements of which it is com-

pounded, and again reconstructing the original metal from these materials. Still less has he realised the dream of the alchemists by transmuting one metal into another. But if we are on this account to refuse to his hypothesis that fair examination to which it is so well entitled, we are establishing a dangerous precedent, and incurring the charge of gross inconsistency. Who, for instance, has seen and handled an atom, or even a molecule? Yet we admit the existence of such portions of matter, we ascribe to them certain properties, and we even speculate on their relative positions in complex bodies, because by so doing we can, to the best of our knowledge, most satisfactorily explain certain phenomena. This is precisely the evidence in favour of the undulatory theory of light or of the kinetic theory of gases. They are accepted as best accounting for the facts of the respective cases. Just so with Mr. Lockyer's hypothesis of the dissociation of the elements; it explains, with a completeness hitherto unattained, a number of difficulties in solar and stellar physics. If we admit that "the solar phenomena in their totality are due to dissociation at the photospheric level, and re-association at higher levels," we can then understand the ascending and descending vertical currents in the solar atmosphere, the absorption in sun-spots, and their association with the faculæ, as well as the seemingly continuous spectrum of the corona and its peculiar structure, the excess of supposed calcium in connection with minimum sun-spots, and, contrarywise, the connection between an excess of so-called hydrogen with a maximum of sun-spots.

Even at present, therefore, until some fact is shown to be irreconcilable with Mr. Lockyer's views, we consider ourselves perfectly justified in giving them our provisional adhesion, as a working hypothesis to be constantly tested by reference to observed phenomena.

As regards what may be called laboratory verification, there is a difficulty in the way which at first sight may seem insurmountable.

Let us suppose that we have volatilised and dissociated an element, say copper, and that we have, floating in the voltaic arc or in any other region of intense artificial heat, the vapours of its unknown constituents, which we may call x and y . The problem is to withdraw these vapours from the arc and from each other without allowing them the opportunity of re-associating and so re-constituting the original copper. If we allow the temperature to decline, this result must, it should appear, inevitably follow. But suppose we volatilise and dissociate two or more elements together; is it not at least conceivable that on re-association they might form new combinations? Thus, suppose we have a vapour mixture of—



Is it not possible that x might combine not with y but with y' or y'' , forming bodies which it would be easy to distinguish from the alloys and other compounds of our reputed elements. It will be of course desirable to submit all our simple bodies to degrees of heat higher if possible than have ever been attained before and under every conceivable modification of circumstances. Above all, spectroscopic research will have to be instituted and scrutinised from this novel point of view. Perhaps by some of these methods we may succeed in penetrating to a deeper, a more fundamental series of elements to which our present supposed simple bodies stand in the relation of mere compounds, and of scrutinising their relations and studying their combinations.

But even if it is not given to man to advance farther than a mere knowledge that our present elements are not ultimate and primary species of matter—and this is all to which Mr. Lockyer, so far, lays claim—it is scarcely possible to over-rate the value of the discovery. Our views of the constitution of the universe are rendered far grander and broader. The atom instead of being, as was

recently thought, a "manufactured article" alike throughout the universe and incapable of change is now seen to depend for its nature and very existence upon the thermic condition of its medium. The elements of the planet are decomposed in the sun to a degree dependent upon its temperature. And perhaps the bodies which there resist all further change, and would to a rational being capable of inhabiting our central orb appear as simple, may be decomposed in hotter stars, or in that great body round which, as some astronomers tell us, all the heavenly bodies are describing a revolution. On the other hand, in planets colder than our earth bodies which we are capable of resolving into components might prove apparently elementary. The "elements" in each planet, comet, and sun are therefore the result of a process not altogether heterologous to the "struggle for existence" which eliminates from our earth all organic forms not in harmony with its general conditions; and from this point of view Mr. Lockyer may be saluted as the Darwin of the inorganic world.

A NEW TEST FOR COBALT.

By T. TATTERSALL.

If a solution of a cobalt salt be mixed with cyanide of potassium, a cinnamon-coloured precipitate takes place, the precipitate, as is well known, being soluble in a slight excess of the reagent to a clear yellow liquid. But if now a few drops of yellow ammonium sulphide be added, a blood-red colour is obtained, even in presence of nickel salt, or any soluble cyanide except the copper compound, which completely prevents the reaction. Colourless ammonium sulphide will not produce the red colour. It may also be obtained, though not with the same degree of delicacy, by using Na_2SO_3 or SnCl_2 .

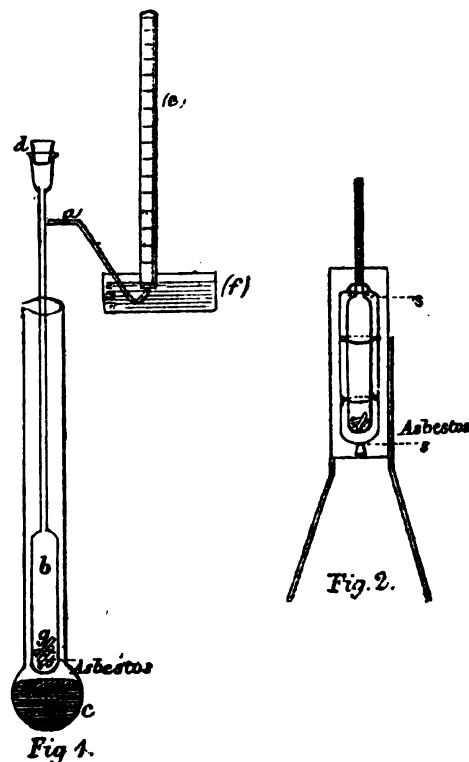
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VICTOR MEYER'S NEW METHOD OF DETERMINING VAPOUR-DENSITIES.

By WATSON SMITH, F.C.S., F.I.C.

I HAD been for some time looking out wistfully for a truly reliable and accurate method, and one not possessing too much intricacy and risks of failure, for the determination of the vapour-densities of the highest boiling hydrocarbons, wherewith I should be able to settle once for all any possible question as to the molecular integrity of the bodies I had considered to be isomeric dinaphthyls. This want has just recently been supplied in the most complete manner, and the point mentioned quite settled, by means of a method so simple, universally applicable, easy and rapid of execution, and withal accurate, that it deserves a foremost place amongst scientific methods. Especially for the worker in the fields of original research will this method indeed be a great boon, and, in fact, amongst methods of its kind, will no doubt take the very first place. Prof. Victor Meyer, of Zürich, has for some time been working upon and perfecting methods of vapour-density determination. The present method is the improved form of one described in the *Deut. Chem. Ges. Ber.*, xi., 1867, and is described by V. Meyer and Carl Meyer in the *Deut. Chem. Ges. Ber.*, xi., 2253 (*i.e.*, the last issue). I have been astonished with the accuracy of the results this method yields, considering the celerity and ease of the determination, and I thought it would be doing good service to help to spread the news of the discovery of a method of vapour-density determination scarcely requiring more time or trouble than the determination of a melting- or boiling-point. With this method the temperature of the bath and the size and contents of the vessel employed do not need to be considered in the calculation.

b (Fig. 1) is the specific gravity vessel of glass, containing 100 c.c., and about 200 m.m. high as regards the cylindrical lower part. The stem above is about 600 m.m. high. d is a caoutchouc stopper closing and fitting well above. a is a narrow, almost capillary, evolution-tube, through which the air escapes in the process, and bubbles up into the graduated tube, e, filled with water, and standing in the little pneumatic trough of glass, f, over water. At the lower part of the apparatus, which must be perfectly clean and dry to begin with, a little asbestos is placed to break the fall of the little glass tube, g, with weighed substance, which at the proper time is dropped in at d. The bath of glass, c, for bodies boiling up to 300°, may be furnished with water, xylene, aniline, ethyl benzoate, amyl benzoate, or diphenylamine, according to the boiling-point of the



substance determined. And these bodies do not need be pure. It is only requisite, in short, to know that the bath furnishes a uniform and constant temperature sufficient easily to vapourise the substance. The substance is weighed out in a little tube, open at one end, and the amount taken should be such that its vapour will occupy somewhat less than half the volume of the cylindrical part of the apparatus, b (fig. 1). The glass bath, c, is placed on wire gauze and heated with a Bunsen's burner. If the substance boil over 300°, then the lead bath (fig. 2) is used. This is merely an iron tube, mounted on three legs. Pieces of lead are thrown in and melted by the heat of a large Bunsen's burner with fifteen jets. To prevent the glass cylinder from touching the iron sides of the tube it is enclosed in a sort of wire cage, as in fig. 2 (s). The apparatus must be cautiously and gradually introduced into the lead bath, else it is cracked or broken. When the lead bath is employed the requisite temperature for vapourising the substance is ascertained simply by placing a little of the substance in a narrow test-tube, and dipping this under the surface of the molten lead, and both observing and applying the ear and listen-

ing if it boils. This is most simply and easily ascertained. The remainder of the process is the same whether lead bath or any other be employed. The instrument, into which a little well-ignited asbestos has been introduced, is gradually and carefully introduced into the bath and fixed in position, the end of its fine tube dipping sufficiently deep below the water in the little trough. The graduated tube stands close by, filled with water and inverted in the trough. The cork, *d*, is now inserted, and it is necessary to wait till the little column of water, extending a m.m. or two up the fine evolution tube, remain quite stationary, no more bubbles and no further motion occurring. This is a sign that a uniform temperature obtains within. The cork is now taken out, and the little tube with substance dropped in, and the cork quickly replaced again, whereby a bubble or two of air escape by displacement. These are not heeded, but immediately afterwards the inverted tube is placed quickly over the point of the evolution tube, *a*. Very soon the substance below is vapourised, and displaces the air of the apparatus above, which issues in a rapid stream of bubbles into the graduated tube. This evolution at an end (generally occupying 10 to 15 seconds), the little water column will first assume a tranquil condition; others gradually show signs of receding. It will recede after some little time a certain distance up the narrow tube, *a*, the effect of diffusion, set up below, between vapour and air. As soon as the gas evolution is over the cork is removed, the thumb placed over the orifice of the graduated tube, and the latter removed and completely immersed in a large deep cylinder filled with water and containing a thermometer. The barometer and thermometer in the water are now, after waiting a minute or two, read off, and then all observations are at an end. The following simple formula gives the vapour density—

$$D = \frac{S \cdot 760(1 + 0.003665t)}{(B - w) V \cdot 0.001293}$$

or gathering together the constants—

$$D = \frac{S(1 + 0.003665t) \cdot 587780}{(B - w) V},$$

where—

S = weight of substance.

t = temperature of water in cylinder, *i.e.*, of the room.

B = Barometric column reduced to 0°.

w = tension of aqueous vapour at *t*°.

V = measured volume of air, in grad. tube.

In the case of substances slightly decomposed by atmospheric air at their boiling and vapourising points, dry nitrogen gas is introduced into the apparatus, and the vapour density is thus determined in nitrogen. *V. Meyer* prepares the nitrogen according to the Gibbs-Böttger method by boiling a solution of 1 part potassium bichromate, 1 part ammonium nitrate, 1 part commercial sodium nitrate, and in 3 parts of water. The gas was also passed over a layer of ignited metallic copper before conducting into the gasholder. From the gasholder it passes first through drying tubes, and then down a long narrow glass tube reaching to the bottom of the vapour density apparatus, itself perfectly clean and dry. The instrument is thus filled by displacement and most simply and expeditiously.

The wide range of this apparatus is shown in the numerous examples of determinations made by Prof. *V. Meyer's* assistant, Herr Carl Meyer, by its means. Chloroform boiling at 61° is first determined (in steam), and, lastly, in the lead bath, the perchlor-diphenyl of Merz, boiling considerably higher than 440°. This determination was made in an atmosphere of nitrogen.

Results.

	Calc.	Found.	Boiling pts.
Chloroform (in steam)	4.13	4.13	61°
Perchlordiphenyl (in nitrogen and lead bath)	17.24	17.43	over 440°

To demonstrate the great value of the method for ascertaining the molecular weights of certain high boiling hydrocarbons, whose compositions may lie so closely together that the centesimal numbers obtained by an organic combustion will fail to distinguish them, it may just be mentioned that the body chrysene, $C_{18}H_{12}$, as regards the numbers furnished by organic analysis, cannot be distinguished from dinaphthyl, $C_{20}H_{14}$. The calculated vapour densities of the two bodies are respectively 7.89 and 8.77. The numbers found by Victor Meyer's method were—for chrysene, 8.12; ditto for dinaphthyl, 8.73. (I am told the substance was not quite pure.)

I am told the apparatus for the above method is to be had on application to Herrn H. Fischli, Zürich, Rami-strasse 45.

Zürich, Feb., 1879.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, February 6, 1879.

Dr. J. H. GLADSTONE, F.R.S., President, in the Chair.

AFTER the announcement of visitors, the minutes of the previous meeting were read and confirmed.

The Treasurer, Dr. RUSSELL, then announced that he had received a bequest of £1000 from the executors of the late Mr. Sidney Ellis, of Nottingham.

The following certificates were read for the first time:—
A. W. Stokes, T. L. Teed, C. H. Hutchinson, G. Rait, W. Stone, T. Griffiths, W. Palmer.

During the meeting the following gentlemen were balloted for and declared duly elected Fellows of the Society:—F. R. Japp, C. F. Cross, H. Wilson, C. E. Cassel, A. E. Menke, J. J. Broadbent, E. A. Letts, E. H. Rennie, W. Stevenson, S. Spencer, C. W. Smith, A. J. G. Lowe, T. Gough, R. Gracey, P. P. Bedson.

The PRESIDENT said that it was scarcely necessary to remind the Fellows that the discussion on Dr. Tidy's paper had been adjourned to this evening, and that it was his duty to preside over the champions of the three processes for determining the organic matter in potable waters, and to see—if he might be allowed the expression—that the lists were clear and the fight was fair. The question to be discussed this evening was that of the relative merit of the processes, and there was the question behind, viz., how far these processes tell us whether a water is fit to drink or not. It was one thing to determine the carbon and nitrogen, and quite another to decide whether such quantities so detected were hurtful or not. He should like to mention that two curves which formed part of Professor Tidy's paper had not been published with the letterpress on account of the delay which had arisen during some corrections. The curves would be published in the forthcoming number. He would call on Mr. Riley, as he moved the adjournment, to open the discussion. Mr. Riley, however, not being present, Dr. Frankland opened the discussion.

Dr. FRANKLAND said that in responding to the President's invitation to open the discussion he wished to express the gratification which the perusal of Dr. Tidy's paper had afforded him. The author had performed his task in a truly philosophical spirit, with fairness and candour. He had frankly told us of the alteration of some of his former opinions, and, above all, he had contributed a large amount of important and laborious work to the subject of water analysis. He thought that chemists generally were much indebted to Dr. Tidy for the clear and impartial style in which he had handled an intricate and exciting theme. As he would have to criticise some parts of the paper

adversely he wished to record at the outset his very favourable opinion of it as a whole. To proceed with the details of the paper, he quite agreed with the author that if it were possible to estimate the organic matter in the water before evaporation, such a mode of estimation would be preferable to any operations of the kind upon a water residue; but unfortunately he knew of no method by which this could be accomplished. He also agreed with the author's opinion that potable waters contained a great variety of organic matters, although he would not go so far as to say that the variety was "infinite." He entirely dissented, however, from the author's suggestion that non-volatile matter may be removed *mechanically* during evaporation. He had proved by actual experiments that mechanical removal only took place during violent agitation or the breaking of gas bubbles at the surface of the liquid. The opinion that solid or liquid particles were mechanically removed during quiet evaporation was emphatically negatived by almost every quantitative determination in analytical chemistry. He was still more astonished, however, to find Dr. Tidy, who is a physician as well as a chemist, suggesting that the poisonous constituents of sewage may be volatile:—"Remembering how virulent that poison is, I should myself be far from astonished if it were of a highly volatile nature, just as we know that arseniuretted hydrogen far exceeds arsenious acid in intensity of action on the human body." All the knowledge we have hitherto acquired about the infectious matter of epidemic disease points to the inevitable conclusion that the propagating material is not merely organic, but organised; and that its virulence, unlike that of arsenic or other similar poisons, resides in its vitality and its power of multiplying itself almost indefinitely in the human body. Now it is impossible to conceive of organisation in substances whose molecules are free to move as they list:—An organised liquid, gas, or vapour, is a physical impossibility, and hence there is no foundation whatever for the opinion that the poison of sewage is volatile, or that it is not contained in the residue left on evaporation. Indeed, the author, somewhat inconsistently, admits this generally-accepted view of the nature of morbid poison when he says, soon after making the foregoing suggestion,—"But it must be remembered that the evils arising from drinking contaminated water are probably the work of invisible particles—germs or whatever else you like to call them." We cannot, however, estimate separately the *disease producing* organic matter of polluted water. It is probably, even in the worst waters, so small in quantity as to be almost imponderable. Dr. Tidy appeared to be equally inconsistent in his great fear of loss of organic matter by oxidation in a *reducing atmosphere of sulphurous acid*, when he afterwards went on to show the extreme difficulty of oxidising such organic matters by an *acidified solution of potassic permanganate*! Moreover, the oxidation of organic matter by no means necessarily increases its volatility. The difficulty which the author has occasionally expressed in the reduction of nitrates when these salts occur in large quantity appears to have arisen from his fear of oxidation by sulphurous acid, and his consequent neglect of the precautions given. He says, "I cannot conceive that the organic matter in the residue can survive direct actual treatment with a sulphurous acid solution. I never do this part of the experiment without a feeling of despair." This conception and this feeling were, however, utterly opposed to facts. Hundreds of combustions of the residues of shallow well-waters containing large quantities of nitrates have demonstrated that the organic matter does survive this treatment. Moreover, this difficulty is in practice more imaginary than real; for when a water contains these excessive quantities of nitrates, it is to be condemned on this ground alone. With regard to the remaining obstacles mentioned in connection with the combustion process, it is easy to procure suitable cupric oxide, either by oxidising sheet-copper in a muffle, or by procuring roasted copper scraps from the makers of sulphate of

copper. The speaker always procured these scraps from Messrs. Tennants, London Road, Manchester, and it was only necessary to heat them to redness in an iron tube for an hour or so in a current of air to fit them for use. There cannot be any occluded nitrogen in the copper, as suggested, because the copper is always heated to redness *in vacuo* before the combustion begins. The author is now convinced of the accuracy of the determination of organic carbon, and the speaker confidently anticipated that further acquaintance with the combustion process would prove to him that the corresponding determination of organic nitrogen was quite accurate enough for all practical purposes. With regard to the albuminoid ammonia process, his experience coincided with that of Dr. Tidy. It had been put forward as capable of answering the question, "Is the water wholesome or is it not?" It does not, however, answer this question: It condemns water containing only peaty matter, and it acquits water containing urea and uric acid! If we look at the results of its application to the waters supplied to London and recorded in this paper, could anybody believe that in 1876, for instance, the Chelsea Company's water, drawn from the Thames, was "safe" in January, "dirty" in March and August, "safe" again in November, and "of extraordinary purity" in December, when it contained no less than 0.423 part of organic elements in 100,000 parts of water; and when three other samples drawn from the same source were denounced as "dirty" by the same process. Again in 1871, whilst the West Middlesex Company's water was "of extraordinary purity" in January, April, and May, the Chelsea Company's water was "dirty" in those months, although quite "safe" in December. Floods undoubtedly make Thames water dirty, but the albuminoid ammonia curve pursues the even tenor of its way quite regardless of them. The very instructive comparative curves, given at the close of the paper, prove conclusively that there is no sort of connection between the pollution of water and the proportion of so-called albuminoid ammonia. He now came to the most important part of the paper, viz., that treating of the oxygen or permanganate process. Experiments had recently been made in his laboratory by Mr. Woodland Toms which showed that there was not, as suggested by the author, any substantial difference between Dr. Tidy's method of performing this process and his own, which was that originally described by the late Dr. W. A. Miller. The author speaks of this process as if by it "everything capable of oxidation" was completely oxidised. But there must be some mistake here, because of the numerous organic matters hitherto submitted to it oxalic acid was the only one completely oxidised. The author had made numerous experiments on the influence of various mineral salts upon the test, and he had shown that the presence of nitrates made no difference with its indications. It was far otherwise, however, with sulphuretted hydrogen, ferrous salts, and nitrites. The first two were easily detected and removed, but the nitrites presented a formidable difficulty, which, in the speaker's opinion, the author had not overcome. He (Dr. Frankland) admitted that they were not often present in potable water; but unfortunately they were especially liable to be present in well-waters, and the oxygen process might thus condemn deep well-waters of most excellent quality. The author's own figures, obtained with the Kent Company's deep well-water, showed this, especially when the very erratic curves of this very uniform water were studied. Thus, in 1870, for instance, the oxygen consumed by this water varied from 0 to 50, whilst the organic elements varied only from 1 to 3.3. In 1871, the oxygen consumed varied from 1 to 7, but the organic carbon and nitrogen only from 1 to 2.6. In 1874, the oxygen consumed was remarkably and uniformly low, whilst the combustion results were, for this water, remarkably and uniformly high. Again, in 1875, the oxygen consumed varied from 1 to 7, but the organic carbon and nitrogen only from 1 to 2; and, lastly, in 1876, the oxygen consumed varied from 1 to 3, but the

organic elements only from 1 to 3. There is a strong probability that the organic matter in the Kent Company's water is acted upon but very slightly by permanganic acid, and that the effects recorded by Dr. Tidy were chiefly produced by nitrites. The speaker had observed on one occasion that a sample of this water, on its first arrival at his laboratory, consumed three times as much oxygen as it did after standing a couple of days, although he had proved that the organic matter in this water underwent no diminution during eleven days' agitation with atmospheric air. Nevertheless there was undoubtedly a remarkable approximation between the curves representing for many years the organic carbon in the river waters supplied to London and those recording the oxygen taken up from potassic permanganate; and the speaker was not without hope that as a short and easy cut to the valuation of organic carbon the process might be made available for potable waters generally, but he thought the author had, as yet, failed to establish it in this position. There were also some other remarks about the capabilities of the oxygen process, which must not pass unnoticed. Dr. Tidy admitted that it certainly does not affect all organic substances; but he says it "affords the most positive evidence (evidence to my mind beyond dispute) of the relative quantity of matters in the water likely to be injurious (for it is the offensive and deleterious we wish to hunt out, and not the inert and the harmless), and enables us to speak with confidence in advising the use of, or the rejection of, a water for drinking purposes." Where is the evidence of this beyond dispute? Dr. Tidy has not given us the slightest fragment of it. If it exists it ought to be produced, because it would put the oxygen process upon the firm and trustworthy basis which it, at present, certainly lacks. No process of water analysis can distinguish between "offensive and deleterious" and "inert and harmless" organic matter; but there is a method which can take account of starch, sugar, gum, gelatine, uric acid, and urea. Dr. Tidy said, and he quite agreed with him, that these substances are in themselves quite innocuous; but how do they get into potable water? People are not in the habit of throwing sugar, starch, gum, or gelatine into rivers. There is a sufficient observance of the proverb "Waste not, want not" to prevent such a use of these valuable materials. Care is taken that these substances first pass through a canal before they reach a river, stream, or well, and that canal is the alimentary canal! Will Dr. Tidy assert that these substances embalmed in the secretions of that canal are *ever* fit for human consumption, or are not, at times, charged with the potentiality of disease? Hence he (Dr. Frankland) said that no process of water analysis is trustworthy which does not take full account of these substances, inert and harmless though they may be, when they leave the grocer's shop. Another statement in the paper in reference to the oxygen process ought not to pass unchallenged, although the author had ceased to act upon it. He said, "In the case of the London waters, however, I think I have abundant experiments to prove that the oxygen used, multiplied by eight, denotes very nearly the actual quantity of organic matter present; nor would it be difficult to obtain further evidence (if it were required) on this point from the work of others." Will Dr. Tidy mention the details of some of these experiments? To chemists generally there is no method known by which "the actual quantity of organic matter present" in London water can be even approximately determined. How, then, can that quantity be proved? If the author knows of a process why does he not disclose it? Such a discovery would go far to sweep the oxygen, albuminoid ammonia, and combustion processes into oblivion, for it could scarcely fail to be equally applicable to other waters besides those consumed in London. With regard to the adjuncts to the oxygen process, the speaker remarked that he considered the colour test to be useful only from a sentimental point of view. Comparatively few organic

matters are coloured, and the coloured part of the organic matter of water is probably a very small fraction of the whole. A water may contain much organic matter, and be nearly, if not quite, colourless, or it may contain but little, and be deeply tinted. Thus, as mentioned by the author, a sample of water which had been in contact with precipitated alumina was, when examined by the speaker, of the very light blue-green tint of *pure* water when seen in a two-foot tube, although it yielded by combustion 0.160 per 100,000 of organic carbon, and another sample which had been in contact with mine mud was, when clarified by subsidence, colourless in a quart decanter, although 100,000 parts of it yielded on combustion no less than 0.544 part of organic carbon. The author, however, said "The peculiar tint of the water is an indication of the kind of organic matter, and the tint-depth is an indication of the quantity of organic matter in the water." The speaker was not aware of the grounds upon which this statement was made. It was not impossible that spectroscopic examination might help us. The absorption-bands of brown urine and of peat might be different, and it was surprising that the author, who evidently attached so much value to the colour test, had not pushed his investigation in this direction. For his own part he had been unable, except in waters from the same district, to guess the approximate portion of peat from the tint-depth of the water. The tinctorial power of the peat in one district seemed to differ from that in another. But it was scarcely necessary to pursue the subject further, as the author himself concludes his advocacy of the 2-ft. tube by virtually condemning it as a quantitative instrument. The quart decanter, or rather flask, was, in the speaker's opinion, more useful than the 2-ft. tube, because if water appears coloured or turbid in such a vessel the consumer would have just cause of complaint, but not otherwise. He cordially agreed with the author's strictures on superficial and rapid analyses of waters, in which the determinations were "chosen with far greater regard to the analyst's time than to the needs of the inquiry." The so-called partial analyses were often very partial, and often brought disgrace upon the profession of chemistry. It was satisfactory to know that the author not only preached but practised this doctrine; the fulness and minuteness of his analyses were well worthy the attention and imitation of chemists. He also agreed with Dr. Tidy in deprecating a hard and fast division of waters into classes, although a suggestive classification was often useful. He would prefer to modify somewhat the division into four classes adopted by the author, making a separate and more liberal scale for upland surface water, which rarely contained anything hurtful, and a somewhat stricter one for spring, well, and river water, the organic matter in which was very liable to come from dangerous sources. He would therefore propose the following classification:—

SECTION I.—UPLAND SURFACE WATER.

Class I.—*Water of Great Organic Purity*.—Containing a proportion of organic carbon not exceeding 0.2 part in 100,000.

Class II.—*Water of Medium Purity*.—Containing from 0.2 to 0.4 part of organic carbon in 100,000.

Class III.—*Water of Doubtful Purity*.—Containing from 0.4 to 0.6 part of organic carbon in 100,000.

Class IV.—*Impure Water*.—Containing upwards of 0.6 part of organic carbon in 100,000.

SECTION II.—WATER OTHER THAN UPLAND SURFACE.

Class I.—*Water of Great Organic Purity*.—Containing a proportion of organic carbon not exceeding 0.1 part in 100,000.

Class II.—*Water of Medium Purity*.—Containing from 0.1 to 0.2 part of organic carbon in 100,000.

Class III.—*Water of Doubtful Purity*.—Containing from 0.2 to 0.4 part of organic carbon in 100,000.

Class IV.—*Impure Water*.—Containing upwards of 0.4 part of organic carbon per 100,000.

He could not admit, however, without further data the translation of these numbers into terms of oxygen consumed. He had studied this translation for several years in connection with the London waters, and had plotted comparative curves from month to month like those exhibited by Dr. Tidy; and his pupil, Mr. Woodland Toms, had also extended the comparison to other classes of waters. From a large number of these results he selected the following to illustrate the difficulties yet to be overcome before a trustworthy translation of oxygen consumed into organic carbon could be made. In the following tables the column headed "Oxygen" contained the amount of oxygen consumed by 100,000 parts of each water, whilst that headed " $\frac{C}{O}$ " gave the multiplier, by which the oxygen consumed was converted into the organic carbon actually found by combustion of the water residue. The amount of carbon thus found was given in the last column, headed "Org. C."

I. London River Water.

Thames.	Oxygen.	$\frac{C}{O}$	Org. C.
Lambeth	0.164	$\times 1.65$	= 0.271
Grand Junction	0.130	$\times 1.65$	= 0.216
Southwark	0.179	$\times 1.64$	= 0.295
Lea.			
New River	0.067	$\times 1.55$	= 0.104
East London	0.079	$\times 1.85$	= 0.146

In these waters the translation is easy; in fact the three Thames waters require practically the same multiplier, and even if this multiplier were applied to the oxygen consumed by the East London Company's water it would not lead to very serious error. Hence the striking agreement in the comparative curves. The case is very different, however, with other classes of waters, and the multiplier has to be greatly altered, as, for instance, in the following tables:—

II. Well-Waters.

Deep Wells.	Oxygen.	$\frac{C}{O}$	Org. C.
Kent Co.'s	0.015	$\times 5.10$	= 0.077
Colne Valley Co.'s ..	0.0135	$\times 6.90$	= 0.094
Hucknall Tokard ..	0.103	$\times 3.60$	= 0.372
Shallow Well.			
Honiton	0.179	$\times 1.80$	= 0.317
Sutton	0.028	$\times 4.00$	= 0.117

III. Sewage.

Lyme Brook + Sewage	0.452	$\times 5.58$	= 2.527
Effluent water from Crewe Sewage Works	0.245	$\times 4.50$	= 1.104

Here the multipliers not only differ for the most part widely from that of Thames water, but what is much more to be regretted they differ enormously amongst themselves. The gross blunders which might be made by using the oxygen instead of the combustion process are, however, most strikingly illustrated by the following table of contrasts, in which each of two pairs of samples requiring approximately the same amount of oxygen are shown to contain enormously different proportions of organic carbon, and require therefore very different multipliers:—

IV. Contrasts.

	Oxygen.	$\frac{C}{O}$	Org. C.
{ The River Marchnant	0.237	$\times 0.57$	= 0.136
{ Crewe Sewage Effluent	0.245	$\times 4.50$	= 1.104
{ Stream at Hinckley..	0.147	$\times 2.96$	= 0.435
{ River Vyrnwy	0.152	$\times 0.89$	= 0.139

But the most hopeless cases are presented in the following table, in which the results yielded by samples taken from the same upland rivers at different dates are contrasted:—

V. Same Rivers at Different Dates.

	Oxygen.	$\frac{C}{O}$	Org. C.
The Vyrnwy, Sep. 4..	0.422	$\times 0.338$	= 0.148
" " " 30..	0.259	$\times 1.03$	= 0.267
" " Dec. 16..	0.041	$\times 2.92$	= 0.120
The Marchnant, Sep. 4	0.286	$\times 0.55$	= 0.156
" " Oct. 10	0.164	$\times 1.46$	= 0.240

The speaker feared that the utmost attainable in this direction would be a translation of oxygen consumed into organic carbon by the use of a separate factor for each description of water. In the meantime, however, the 1418 cases of approximate coincidence out of 1686 mentioned by the author were very encouraging; and to all chemists who shirk the labour of combustion he strongly recommended the oxygen process interpreted by Dr. Tidy as one far superior to any other, and one which, although rough and ready, would probably only rarely lead them far astray. But in all cases of moment and importance, where the health of large communities was at stake, reliance ought never to be placed upon a method which deals only with the carbon and hydrogen of the organic matter and leaves the nitrogen unnoticed. In conclusion, he congratulated the author on his valuable addition to the literature of water analysis, and on the clearness with which he had brought a great array of facts before the Society. His comparative tables were of great value, and he trusted they would be carefully studied by all chemists who practise water analysis. They formed a chart upon which the rocks so fatal to many a water analysis were so clearly shown that he who runs might read.

At the conclusion of Dr. Frankland's address a general wish was expressed to hear Mr. WANKLYN, who made the following remarks. Unfortunately, he was not able to take so favourable a view of the paper as Dr. Frankland. He would restrict himself to the subject of the paper, and hardly say anything about the paper itself. He believed that he was not bound to notice the paper, and should therefore deal with the subject of it. One of the subjects was the controversy between Dr. Frankland and himself, which commenced in 1867 in the pages of a journal no longer published—*The Laboratory*—and as long ago as that date Chapman, Smith and himself directed attention to the cardinal defect of the combustion process, and this defect had never been overcome. It is this, that the organic matter in the water does not survive the evaporation to dryness. This may be doubted; but let us consider what is the organic matter. We have modified cellulose which is very sensitive to the action of dilute acids; e.g., in a few seconds a solution of cane-sugar, when treated with dilute sulphuric acid, becomes converted into grape-sugar, so as to reduce the copper test, once we get cellulose into the plastic condition it does not need oxidation to break it up. If the organic matter of the residue is not that of the water, it is plain that the process is not valid, and this consideration and the practical difficulty of the process have prevented Mr. Wanklyn from making many water analyses by the combustion method. Dr. Frankland said that he would be content if he had a process to burn up the organic matter in the water itself. Messrs. Cooper and Wanklyn have invented such a process, and have burned up substances in solution. The results obtained are interesting. The actual organic matter present in the London Companies' waters does not exceed 2½ milligrams per litre. Dr. Frankland has found 3.99 milligrams per litre as a mean of many experiments: in his paper, the quantity really does not exceed 2½. After this discrepancy comment on the process is needless.

Mr. KINGZETT said that with regard to the occlusion of nitrogen by copper, Thudichum and himself had shown that copper prepared from the nitrate is quite free from nitrogen. Prof. Tidy's experiments on page 78 proved that permanganate practically did not act on gelatin. As

albuminoids were admitted to be dangerous by all, this fact was a stumbling-block in the use of the process, as it overlooked albuminoids.

Prof. BISCHOF said that Dr. FRANKLAND's reply had been so exhaustive that he would confine himself to a few remarks. Though all agreed that it would be preferable to estimate the organic matter in the water, yet we must recollect that by evaporating the water the organic matter was concentrated, and so could be estimated with even more accuracy than in the water itself. He asked Prof. Tidy whether he had made any experiments by condensing the vapour given off, during the evaporation of water for analysis, which proved that a loss of organic matter had taken place, so as to be able to say the oxygen process has detected so much volatile organic matter in the distillate? His plan of evaporating water prevented to a great extent the destruction of organic matter, because it avoided the frequent alternate exposures to hot water and air. Could Prof. Tidy detect by the oxygen process any difference between milk infected, say, with typhoid and scarlet fever and the same milk not so infected? He would like to know the precise meaning attached by Prof. Tidy, in his reports of the London waters, to the words "efficiently filtered." Also, why Prof. Tidy continues to use the ammonia method in his reports to the Medical Officers of Health when he has admitted that the results of that method are unsatisfactory. In conclusion, he would submit to the President the figures obtained by Prof. Tidy and Dr. Whitmore, both using the albuminoid ammonia method, in their analyses of the West Middlesex and Grand Junction waters, figures which differed by nearly 100 per cent, and asked if Mr. Wanklyn or anyone else could account for this discrepancy.

Mr. WANKLYN remarked that he had never had any difficulty in obtaining his alkaline permanganate free from ammonia, and could not tell what results might be arrived at by using impure solutions of permanganate, and, allowing for the impurity: it was also very important that chemists should be trained in a correct manner.

Dr. VOELCKER said that an impartial observer would come to the conclusion that all methods were more or less defective, and some gave very erroneous results. He would most earnestly urge the importance of determining all the constituents of a water, organic and inorganic, and not founding our opinion on one factor.

Mr. GROSJEAN said that all must have recognised the great advantage of having a paper printed before it was discussed, and trusted that it would form a precedent and ultimately the rule. Prof. Tidy did not seem to have standardised his permanganate, but had apparently assumed it to be pure. It was usually standardised, and hence probably the origin of the multiplication of the oxygen used by 8, as the equivalent of oxygen $\times 8$ equals nearly the equivalent of oxalic acid. The commencement of the action of permanganate on oxalic acid is characterised by a tardiness which was explained by Vernon Harcourt, *Chem. Soc. Journ.*, 1867.

Dr. DUPRE protested against the assumption that a chemist who used the albuminoid ammonia process did so to shirk the trouble of the combustion process; and, moreover, did not see why the time, expense, and trouble of a process should not have a certain weight. He had used both the ammonia and permanganate processes, and found the results fairly concordant, with, however, strange exceptions now and then. He had not had any difficulty in preparing alkaline permanganate solution free from ammonia. He gave as an instance of concordant results obtained by the ammonia process, the results obtained by six chemists in examining 249 waters:—Of these, 66 were first class waters, and all agreed that 64 were first class waters and 2 second class; 60 were second class waters, and all agreed that 53 were second class, 6 first class, 1 third; 123 were cistern waters, and 6 were put in the first class, 81 in the second, and 36 in the third.

Mr. W. THORP asked Prof. Tidy what his conception was as to the nature of the organic impurity, as he had

compared it to hydrocyanic acid, &c. He did not think that any free sulphuric acid was likely to be formed during the evaporation, as the carbonates present would be more than sufficient to neutralise any that might be formed, and if there was any chance of such an occurrence it might be avoided by the addition of a little pure sulphite of sodium. He did not think that Prof. Tidy had done justice to Nesslerising. When in work his (Mr. Thorp's) error in estimating tints was only 3 per cent. Railway men were not accustomed to judge minute differences, and so their arrangement of Nessler tints was by no means a fair experiment. He thought great difficulty would be experienced in applying the oxygen process to waters containing nitrites. Moreover, some waters containing no nitrites rapidly reduced permanganate. He did not place much reliance on the colour of a water.

Dr. HAKE remarked that when copper was cooled in CO_2 no occlusion took place.

Prof. TIDY thanked the Fellows for the kind manner in which they had received his paper, and expressed his satisfaction at having provoked such an interesting discussion. He had concluded that the poison in water was volatile from analogy, because when it did act it was very virulent, and virulency and volatility usually go together. He relied on the colour of a water more as a useful adjunct to the results obtained by analysis. In the Thames water the oxygen used $\times 8$ gives the difference very nearly between the inorganic salts and the total residue.

The warmest thanks of a crowded meeting were carried by acclamation and tendered to Prof. Tidy for his paper.

The Society then adjourned to February 20, when the following papers will be read:—"Investigations into the Action of Substances in the Nascent and Occluded Conditions (Hydrogen, continued)," by Dr. Gladstone and Mr. Tribe; "On some Methods of Vapour-Density Determination," by J. V. Brown; "On the Quantitative Blowpipe Assay of Mercury," by G. Attwood.

PHYSICAL SOCIETY.

Annual Meeting, February 8, 1879.

Prof. W. G. ADAMS, President, in the Chair.

THE PRESIDENT read the report of the Council, which showed that the papers had been more numerous during the past than in any previous year, and that their value and interest had been well sustained. A copy of the collected papers of the late Sir Charles Wheatstone was laid on the table, and the work will shortly be issued to the members of the Society. The President then gave a brief review of the physical work of the past year, dwelling more especially on the papers read at the meetings.

Votes of thanks were then passed to the President, to the Lords of the Committee of Council on Education, to the Demonstrator, Treasurer, Secretary, and Auditors, and the following were elected as Council and Officers for the ensuing year:—

President—Prof. W. G. Adams.

Vice-Presidents—Prof. G. C. Foster, Prof. R. B. Clifton, Lord Rayleigh, Dr. Spottiswoode, Sir W. Thomson.

Secretaries—Prof. A. W. Reinold and Mr. W. Chandler Roberts.

Treasurer—Dr. E. Atkinson.

Demonstrator—Prof. F. Guthrie.

Other Members of Council—Capt. W. de W. Abney, Dr. Warren de la Rue, Major E. R. Festing, Prof. Fuller, D. Huggins, Prof. A. B. W. Kennedy, Prof. McLeod, The Earl of Rosse, Mr. G. Johnstone Stoney, Dr. Wormell.

Honorary Members—Prof. G. R. Kirchhoff, Dr. J. Plateau.

The meeting was then resolved into an ordinary one, and Dr. O. J. LODGE read a short paper on a method of calculating the curve of temperature in a rod along which heat is being conducted.

Mr. SHOOLBRED gave an account of electric lighting, illustrated by diagrams of the most recent magneto- and dynamo-electric machines and examples of the lamps in vogue. The only surviving magneto-machine is that of De Meriten's, which is incomparably superior to the older ones of Nollet and Holmes. The dynamo-electric machines described were the continuous-current machines of Siemens, Gramme, Wallace-Farmer, and the alternating-current machines of Wilde, Gramme, and Lontin. Wilde's machine is the first of these, or parent machine, and Lontin's so resembles it that the latter cannot be used in England. In these machines the current from a continuous machine is passed through a second machine, which yields the alternating currents. In Lontin's machine, also, a number of distinct currents are generated in separate circuits, each of which is capable of feeding several lights. There is now one in use on the Western Railway of France which gives three distinct currents, each of which supplies four different lamps, making a total of twelve lights. The American Brush machine was also mentioned. The Dubosq lamp, which was the first regulator, is well adapted for laboratory purposes, but for practical purposes the Serrin is preferable. Rapiéff's lamp is used in the *Times* office. The De Mersanne, which was highly spoken of at the Paris Exhibition, moves the carbons by bevelled gearings. The Wallace-Farmer lamp, though durable, is unsteady, perhaps because only inferior gas carbon has yet been used. Jablochkoff's candle was found to be defective from the solid insulator, such as plaster, used between the carbon. This made it very expensive also. Experiments in Paris had shown that whereas Jablochkoff's system cost 10d. per hour per light, the other systems cost only one half of that. In Wilde's candle the solid insulator was dispensed with, air taking its place, the arc always tending to keep at the top of the candle by electro-dynamic repulsion. In the De Meriten's candle three strips of carbon were used, the intermediate one being a stepping-stone to the arc which passes between the two outer ones. Werdermann's and Reynier's so-called incandescent lamps were also shown. Mr. Shoolbred, after alluding to the fact that the upper (positive) carbon takes a crater form, and hence becomes a reflector shedding the light downwards, stated that experiments had proved the line of maximum intensity of light to pass downward at an angle of 60° to the axis of the vertical carbons. By giving the positive carbon a horizontal displacement behind the lower negative one, Mr. Douglas, of the Trinity House, had been able to raise this line till it became horizontal, an advantage in light-houses. He also pointed out that, whereas in Paris the

Jablochkoff waxed for a period short compared to that in which it waned, in London it waxed for longer than it waned, which was of course an improvement; and Mr. Shoolbred suggested that it might be due to the fact that the engine worked at a speed nearer to that of the machine, and that the machine was founded more solidly in London than in Paris.

Mr. WERDERMAN said that it was a mistake to call his lamp an incandescent one, the fact being that all carbon lamps gave light from the incandescence of the positive carbon, and that a small arc was formed in his lamp between the two electrodes, which could be varied by the pressure between them. He maintained that it was as easy to produce 500 lights as 10 from the electric light by sub-division, as he hoped soon to show, and stated that the size of the carbons greatly controlled the intensity of the light.

Prof. AYRTON held that the obstacle to the sub-division of the electric light was not an electrical one, but was due to the fact that the amount of light produced by the current is not in direct proportion to the amount of the heat produced.

In contradiction to Prof. Ayrton, Mr. WERDERMAN stated that in the electric arc the opposing electromotive force was proportional to the original electromotive force.

Prof. SYLVANUS P. THOMSON pointed out that residual magnetism in the cores of the bobbins of dynamo-electric machines lowered their efficiency, and hence short cores, as in the Wallace-Farmer machine, were an improvement.

EDINBURGH UNIVERSITY CHEMICAL SOCIETY

January 29, 1879.

ALEXANDER MACFARLANE, D.Sc., F.R.S.E., in the Chair.

A PAPER was read by Mr. J. S. THOMSON, on "*Paraffin and what is got from it, as illustrated by Exhibits at the Paris Exhibition of 1878.*" A full account was given of the modes of preparation of these exhibits, which consisted of specimens of all the commercial products of the Addiewell Chemical Works. These included naphtha, burning oils, lubricating oils, solid paraffin in blocks (weighing not less than 7 cwts. each), sulphate of ammonia, and candles of various kinds made from the paraffin wax. The methods of preparation of other products of paraffin were also explained; of normal paraffins, alkaline bases of the leucoline series found in the acid tars, phenols and analogous bodies found in the soda tars, chrysin and paraffins of various melting-points, from 21° to 65.5° C.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

JANUARY, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

	Appearance in 2 foot Tube.	Ammonia.		Nitrogen as Nitrate, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia.	Chlorine.	Sulphuric Anhydride.	Hardness on Clark's Scale.	
		Saline.	Organic.								Before Boiling.	After Boiling.
		Gr.	Gr.	Gr.	Gr.	Gr.	Gr.	Gr.	Gr.	Gr.	Degs.	Degs.
<i>Thames Water Companies.</i>												
Grand Junction	Slightly turbid	0.000	0.012	0.165	0.130	21.40	9.230	0.720	0.72	1.600	13.2	5.10
West Middlesex	Clear	0.000	0.015	0.165	0.120	20.00	9.120	0.648	0.72	1.330	14.3	4.60
Southwark and Vauxhall	Slightly turbid	0.000	0.011	0.105	0.135	18.20	6.770	0.756	0.72	1.230	11.5	5.10
Chelsea	Clear	0.000	0.007	0.210	0.033	20.10	9.010	0.648	0.72	1.530	13.2	3.30
Lambeth	Clear	0.000	0.009	0.135	0.037	21.30	7.590	0.612	0.72	1.100	13.7	3.70
<i>Other Companies.</i>												
Kent	Clear	0.000	0.003	0.435	0.002	29.30	10.580	0.936	1.29	3.260	19.4	6.50
New River	Clear	0.000	0.007	0.165	0.030	21.30	8.730	0.684	0.72	1.200	14.3	3.70
East London	Clear	0.000	0.011	0.135	0.049	23.00	7.630	0.720	0.79	1.160	14.3	3.70

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 3, January 20, 1879.

Observations on M. Pasteur's Second Reply.—M. Berthelot.—The author points out that M. Pasteur, though declaring beer to be anærobian, admits that this supposition has not been demonstrated.

Reply to the Notes of M. Trécul of Dec. 30 and Jan. 13.—M. Pasteur.—A continuation of the controversy on anærobian and anærobian organisms.

Temporary Magnetic Powers Developed by Influence in Certain Specimens of Nickel and Cobalt as Compared with Iron.—Henri Becquerel.—Bars of nickel become saturated more rapidly than bars of iron, and consequently setting out from a certain intensity a bar of nickel becomes a temporary magnet almost constant, while iron becomes magnetised more for growing intensities. The difference between the magnetic properties of bars of iron and nickel are so much the more considerable as the conditions of magnetic are nearer to those which determine saturation in one or the other.

Classification of Colours and the Means of Reproducing Coloured Appearances by Three Special Photographic Proofs.—C. Gros.—The author distinguishes two categories comprised under the word colours: lights and pigments. The elementary lights which by their mixture produce all kinds of shades are the green, violet, and orange rays. The elementary pigments which by their mixture produce all kinds of shades are red, yellow, and blue. To obtain directly the elementary tints of rays and of pigments it is sufficient to look through a prism at a white stripe upon a black ground, and at a black stripe upon a white ground. In the first case we see an orange, green, and violet spectrum, and in the latter case a blue, red, and yellow spectrum. In the former case the orange, green, and violet are elementary rays, and in the latter the red, blue, and yellow are rays combined two and two. The author then exhibits and describes an apparatus, which he names the chromometer, and by means of which he produces the protographic effect above mentioned.

Researches on the Effects of Induction Across Telephonic Circuits by Means of the Microphone and the Telephone.—D. Hughes.—Taken from an English source.

New Voltaic Element with a Constant Current.—A. Héraud.—The author uses as an exciting liquid hydrochlorate of ammonia; the depolarising body is mercurous chloride. When the circuit is closed the hydrochlorate of ammonia in presence of zinc gives zinc chloride, forming ammonia and hydrogen, which two bodies go to the positive electrode. The hydrogen reduces the mercurous chloride, producing metallic mercury, hydrochloric acid, and consequently hydrochlorate of ammonia. As long as there exists mercurous chloride around the positive electrode, hydrochlorate of ammonia will be regenerated.

Tetric Acid and its Homologues.—Eug. Demargay.—An account of the tetric, pentic, hexic, isohexic, and heptic acids. These acids agree very closely in their chemical properties, and are coloured a violet-red by ferrous chloride.

Reimann's Fürber Zeitung,
No. 47, 1878.

Dr. Lunge confirms the statements of Miller on the use of tropeolin OO for the titration of soda, for deter-

mining the free acid in sulphate of alumina, &c. So long as no free acid is present the yellow colouring-matter remains unchanged. If there is the smallest excess of hydrochloric acid the colour turns first to magenta, then to orange, and after a few seconds disappears entirely. The author has tested a number of the azo colouring-matters, and has found that they are all indifferent to free carbonic acid, and are capable of marked changes of colour in presence of the slightest excess of mineral acid, or inversely. These changes are most beautifully shown by Poirrier's Orange III. (dimethyl-anilin-diazosulphobenzolic acid), by diazo- α -sulpho-naphthyllic acid, and amido-benzol. Tropeolin OOO is suitable for the detection of alkali (caustic or carbonated), as it undergoes the opposite change of colour from tropeolin OO, being yellow in an acid solution and magenta-red in alkaline liquids. Weaker acids—such as the sulphurous, oxalic, &c.—produce less distinctly marked changes of colour. Acetic acid, like the carbonic, is indifferent to the azo-compounds. Hyposulphite of soda behaves exactly like the salts of strong mineral acid. With the aid of these indicators soda-ash, &c., may be titrated in the cold.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 16, December 19, 1878.

This issue contains no original chemical matter.

No. 17, December 26, 1878.

The only original chemical matter in this issue is a description of the celebrated aniline-colour works of A. Poirrier, at St. Denis, and of the dyes which have been either invented or at least manufactured there. The writer enumerates as such—Violet (produced with potassium bichromate); dimethyl-anilin violet, otherwise known as Paris violet; benzylated dimethyl-anilin violet, purer in tone than Paris violet, and capable of being applied upon wool in presence of acids; methyl-anilin green, formed by acting upon the violet with methyl-chloride; dibenzyl-anilin green, which has not proved commercially successful by reason of its very sparing solubility; phenylen-diamin brown and chrysoidin; cachou de Laval, otherwise known as the patent colours of Croissant and Bretonnière, formed by the action of sodium sulphide upon organic matter. (This colour does not seem to possess any distinctive novelty, and, though exceedingly fast, yields dull shades.) Mention is next made of the oranges and rocellin. The last of these colours is obtained by the reaction of the diazoic derivative of sulpho-conjugated naphthylamin upon naphthol B; the shade produced is very like that of orchil, but brighter, faster, and more economical.

Nos. 1 and 2, January 2 and 9, 1879.

These issues contain no original chemical matter.

Revue Universelle des Mines, de la Metallurgie, &c.,
Tome 4, No. 2, September and October, 1878.

The only chemical paper here is a note by Prof. J. B. Caldwell on the determination of phosphorus in ores of iron, and in cast-iron and steel, taken from the *New York Metallurgical Review*.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 14, 1879.

Transformation of Dibromethylen into a Keton with Four Atoms of Carbon by Means of Hypobromous Acid.—E. Demole.—The author considers it as established that on the oxidation of dibromethylen there is really formed a transition compound, bromacetylen, in the nascent state. At low temperatures this compound

becomes polymerised, resuming HBr, whilst at higher temperatures it takes up simultaneously oxygen and hydrobromic acid, and is converted into bromacetyl-bromide.

Separation of Zinc and Nickel.—F. Beilstein.—The sufficiently diluted solution of the nitrates or sulphates is mixed with ammonia till an alkaline reaction is obtained, and then acidified with pure citric acid. When the solution is perfectly cold sulphuretted hydrogen is introduced till the liquid has a distinct smell, which is generally effected in five to ten minutes. If much zinc is present sulphuretted hydrogen is introduced for five minutes at a time, letting the liquid stand after each introduction, and repeating this till the smell of the gas does not disappear on standing. In this manner we avoid a needlessly long treatment with sulphuretted hydrogen by which traces of nickel sulphide may be carried down. The precipitate of zinc sulphide is allowed to stand for twenty-four hours in the cold, and is then weighed as such. The filtrate is evaporated to a small volume, and after supersaturation with ammonia the nickel is thrown down electrolytically, in which operation care must be taken that no sal-ammoniac is present, as it hinders the precipitation of nickel. The solution should be a nitrate.

On β -Chlorocymol from Thymol.—E. v. Geichten.—A hypothetical and controversial paper, not susceptible of useful abstraction.

Certain Fluorine Compounds of Vanadium.—H. Baker.—A valuable and detailed account of the formation and properties of the fluoxy-vanadates of potassium, ammonium, and zinc, and of fluoxy-hypo-vanadate of ammonium.

Certain New Derivatives of Benzoic Acid.—P. Griess.—The author describes ϵ -nitramido-benzoic acid, ϵ -oxy-nitro-benzoic acid, α -oxy-nitro-benzoic acid, and γ -oxy-nitro-benzoic acid.

Halogen Substitution-products of Ethan and Ethylen (Eleventh Communication).—W. Stædel.—An account of the hexa-chloride of carbon and of the somewhat numerous chlorobromo- and bromo-substitution products of ethan and ethylen.

Compounds of Organic Bases with Mercuric Chloride.—Otto Klein.—In this memoir the author describes certain mercuric compounds of dimethyl-anilin.

On Isoindol (Thirteenth Communication).—W. Stædel and M. Kleinschmidt.—Isoindol presents one of the most interesting cases of pleochroism. If a columnar crystal is held to the light and slowly turned on its axis it appears in succession green, yellow, deep red, blue, and indigo-blue.

Dinitro-benzo-phenon and Dioxy-benzo-phenon.—W. Stædel and E. Sauer.—Dinitro-benzo-phenon is most readily obtained by the oxidation of dinitro-diphenyl-methan. Dioxy-benzo-phenon is best prepared from the hydrochlorate of diamido-benzo-phenon.

Brom-nitro- and Brom-amido-anisols.—W. Stædel and G. Damm.—Not suitable for abstraction.

On Uvic Acid.—W. Stædel.—During an attempt to resolve synthetic uvic acid into dextro- and lævo-tartaric acid a solution of the neutral sodium-ammonium salt was prepared. From this solution there were at first deposited large well-developed crystals of the monoclinic system, on which there were no hemihedral surfaces, and whose solution was optically inactive. In the mother-liquor of these crystals there appeared the rhombic crystals of the two sodium-ammonium tartrates, with their usual hemihedral surfaces.

On Diphenyl Bases.—H. Schmidt and G. Schultz.—The authors have undertaken a complete examination of all the known amido-compounds of diphenyl.

Supplementary Remarks on Alstonia Constricta.—O. Hesse.—The author, in the former paper, questioned the presence of quinin in Alstonia bark. He now learns

that Baron Muller, of Melbourne, who discovered the species, also denies that it contains quinin.

On Diphenyl Benzols.—H. Schmidt and G. Schultz.—The authors seek to ascertain the relation—if any—of violanilin to diphenyl-benzol.

Amidins and Thiamids of Mono-basic Organic Acids.—A. Bernthsen and H. Trompeter.—The authors give an account of ethenyl-tolyl-amidin, benzenyl-naphthyl-amidin, ethenyl-naphthyl-amidin and certain of its salts, benzo-thio-toluidid, aceto-thio-toluidid, benzo-naphthyl-thiamid, aceto-naphthyl-thiamid, benzamido-tolyl-thiamid, benzyl-anilin, and ethyl-naphthyl-amin.

Certain Double Salts of Bivalent Iridium.—C. Seubert.—An examination of the iridic double salts of sodic bisulphite.

Atomic Weight of Iridium.—C. Seubert.—The author's determinations give as a mean result 192.744 ($H = 1$). He points out that Berzelius admits the presence of osmium in the iridium which he employed.

On Absorption Spectra.—J. Landauer.—A spectroscopic examination of safranin. The author not merely agrees with Vogel in doubting the universal applicability of the proposition that every chemical compound has its specific spectrum, but concludes that absorption spectra throw light upon the constituents of a compound only in so far as the colour of a body can be regarded as characteristic of its chemical composition. For the accurate determination of colours the spectroscope is as serviceable as is the microscope for the determination of forms.

Action of Dry Gaseous Hydrochloric Acid upon Sulphates.—C. Hensgen.—In this memoir the author describes the action of dry hydrochloric acid gas upon ferrous sulphate. At elevated temperatures the anhydrous salt is decomposed, and pure anhydrous ferric chloride is formed. From the hepta-hydrated salt was obtained $Fe_2Cl_6 \cdot 2H_2O$ in green acicular crystals, and $FeSO_4 \cdot 6H_2O$ in tabular plates.

MISCELLANEOUS.

Royal Commission.—The Queen has approved the appointment of Mr. Warrington W. Smyth, F.R.S., Sir George Elliot, M.P., Mr. F. A. Abel, C.B., Mr. Thomas Burt, M.P., Mr. Robert Bellamy Clifton, F.R.S., Prof. Tyndall, F.R.S., Mr. Lindsay Wood, and Mr. William Thomas Lewis, as Her Majesty's Commissioners for the purpose of inquiring and reporting whether, with respect to the influence of fluctuations of atmospheric pressure upon the issue of fire-damp from coal, to the adoption and efficient application of trustworthy indicators of the presence of fire-damp, and generally to systematic observation of the air in mines, to improved methods of ventilation and illumination, to the employment of explosive agents in the getting of minerals, and to other particulars relating to mines and mining operations, the resources of science furnish any practicable expedients that are not now in use and are calculated to prevent the occurrence of accidents or limit their disastrous consequences.—*Times*.

Moist Water Colours for Students.—Lech tier Barbe, and Co. have issued a set of student's moist water colours in tin pans. These moist colours are said to be pure, unadulterated pigments. The use of water colours in the moist state, with artists, has almost superseded that of dry cake colours. The latter may have a superiority in some instances, yet the moist, affording a supply of colour with more freedom and rapidity, have come to be generally preferred, and the only obstacle to their being more extensively used still has hitherto been their high price. The shilling box contains the ten colours and three brushes as recommended by the Society of Arts. The main and novel feature of the box is that the colours are

contained in small tin pans; that these pans are removable, and that, when used up, any of them can be replaced by another one, just as in artists' colour boxes. Thirty of the most useful moist colours are manufactured and sold in those small tin pans; so that additional colours may be put in the box, in order to bring it fully within any object in view in water-colour drawing.

MEETINGS FOR THE WEEK.

- MONDAY, 17th.**—Medical, 8.30.
London Institution, 5.
Society of Arts, 8. "Dwelling Houses: Their Sanitary Construction and Arrangements," by Dr. W. H. Corfield, M.A. (Cantor Lectures)
- TUESDAY, 18th.**—Civil Engineers, 8.
Royal Institution, 3. "Animal Development," Prof. Schäfer.
Zoological 8.30.
- WEDNESDAY, 19th.**—Society of Arts, 8. "Turkish Resources and their Ready Development," by J. L. Haddan.
Meteorological, 7.
Society of Public Analysts, 8. "On the Influence of the Decomposition in Butters from Age on the Specific Gravity of the Fat and the Percentage of Soluble and Insoluble Acids," by E. W. T. Jones, F.C.S. "Notes on the Analysis of Butter," by J. M. Milne, M.D. "On Condensed Milk," by O. Hehner, F.O.S. "On Analysis of Coffee Leaves," by O. Hehner, F.C.S. "On the Falsification of Milk," by C. A. Cameron, M.D.
- THURSDAY, 20th.**—Royal, 8.30.
Royal Institution, 3. "Sound," Prof. Tyndall.
Royal Society Club, 6.30.
Chemical, 8. "Investigations into the Action of Substances in the Nascent and Occluded Conditions: Hydrogen (continued), by Dr. Gladstone and Mr. Tribe. "On some Methods of Vapour Density Determination," J. T. Brown. "On the Quantitative Blowpipe Assay of Mercury," by G. Attwood.
- FRIDAY, 21st.**—Royal Institution, 9. "A New Chemical Industry," Prof. Roscoe.
- SATURDAY, 22nd.**—Royal Institution, 3. "Lessing," by Reginald W. Macan
Physical, 3. "On a Current Regulator," by Dr. C. W. Siemens, F.R.S. "On a New Theory of Terrestrial Magnetism," by Profs. Ayrton and Perry. "On the Spectrum of Lightning," by Dr. A. Schuster.

TO CORRESPONDENTS.

By an oversight the source of the article by Profs. Thomson and Houston in our last number was omitted. It was taken from the *Journal of the Franklin Institute*.

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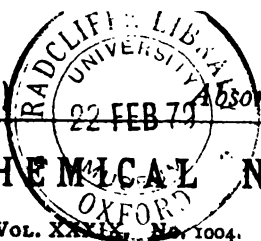
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THE CHEMICAL NEWS.

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Absorption of Gases by Charcoal.

77

ABSORPTION OF GASES BY CHARCOAL.

PART II.

ON A NEW SERIES OF EQUIVALENTS OR MOLECULES.*

By R. ANGUS SMITH, Ph.D., F.R.S.

In the *Transactions of the British Association*, 1878, Norwich, on page 64 of the *Abstracts*, there is a preliminary notice of an investigation into the amount of certain gases absorbed by charcoal. I made the inquiry from a belief previously expressed in a paper of which an abstract is in the *Proceedings of the Royal Society*, page 425, for 1863. I said in that paper that the action of the gas and charcoal was on the border line between physics and chemistry, and that chemical phenomena were an extension of the physical; also that the gases were absorbed by charcoal in whole volumes, the exceptions in the numbers being supposed to be mistakes. The results given were:—

Hydrogen	1.00
Oxygen	7.99
Carbonic oxide	6.03
Carbonic acid	22.05
Marsh-gas	10.01
Nitrous oxide	12.90
Sulphurous acid	36.95
Nitrogen	4.27

It was remarked that the number for nitrogen was probably too low; I had some belief that the charcoal retained a certain amount which I had not been able to estimate.

For common air, the number 40.065 crept into the paper or abstract instead of the quotient 7.06

I considered the numbers very remarkable, but was afraid that they would be of little interest unless they could be brought more easily under the eyes of others; my experiments were somewhat laborious; the exact numbers were seldom approached by the single analysis, but were wholly the result of a series of irregular averages and apparently irregular experiments. The cause of this was clear, as I believed, namely, the irregular character of the charcoal with which I had to deal. The experiments were forgotten, I suppose, by most men, but the late Prof. Graham told me that he had repeated them with the same results that I had published. I might have considered this sufficient, but waited for time to make a still more elaborate investigation of the subject, and to take special care with oxygen, in the belief that, the rule being found, the rest of the inquiry would be easy; this was extended to nitrogen, but not by so many experiments as with oxygen. I am now assured of a sound foundation for inquiries, which must take their beginning from the results here given.

It is found that charcoal absorbs gases in definite volumes, the physical action resembling the chemical.

Calling the volume of hydrogen absorbed 1, the volume of oxygen absorbed is 8. That is, whilst hydrogen unites with eight times its weight of oxygen to constitute water, charcoal absorbs eight times more oxygen by volume than it absorbs hydrogen. No relation by volume has been hitherto found the same as the relation by weight.

The specific gravity of oxygen being 16 times greater than hydrogen, charcoal absorbs 8 times 16, or 128 times more oxygen by weight than it does hydrogen. This is equal to the specific gravity of oxygen squared and

divided by two $\frac{16^2}{2}$, or it is the atomic weight and specific gravity multiplied into each other, 16×16 , and divided by two $\frac{256}{2} = 128$.

Nitrogen was expected to act in a similar way, but it refused. The average number of the latest inquiry is 4.52, but the difficulty of removing all the nitrogen from charcoal is great, and I suppose the correct number to be 4.66. Taking this one as the weight absorbed, $14 \times 4.66 = 65.3$, or it is $\frac{14^2}{3}$. Oxygen is a dyad; nitrogen a triad.

We have then carbonic acid not divided, but simply 22 squared = 484.

Time is required for full speculation, but the chemist must be surprised at the following:—

Carbonic oxide	6	volumes.
Carbonic acid, CO ₂	6+16	" = 22.00
Marsh-gas, CH ₄	6+4	" = 10.00
Protoxide of nitrogen, NO	8+4.66 (N) (4.9)	= 12.66

These four results belong to the early group not corroborated lately, but so remarkably carrying out the principle of volume in this union, giving numbers the same as those of weight in chemical union, that they scarcely require to be delayed.

I am not willing to theorise much on the results; it is here sufficient to make a good beginning. We appear to have the formation of a new series of molecules made by squaring our present chemical atoms, and by certain other divisions peculiar to the gases themselves. Or it may be that the larger molecule exists in the free gas, and chemical combination breaks it up. These new and larger molecules may lead us to the understanding of chemical combinations in organic chemistry and whenever there is union not very firm, and may also modify some of our opinions on atomic weights and the motion of gases.

Of course, I cannot pretend to give the result of these results; but as we have here the building up of a molecule by volumes, so as to form an equivalent of physical combination analogous to the chemical equivalent, it is impossible to avoid seeing that it indicates the possibility of our present equivalents being made up in a similar manner.

I did not expect these numbers; but I certainly, as my previous paper showed, had in full view a necessity for some connexion between physical and chemical phenomena more decided than we possessed.

VOLUMETRIC ESTIMATION OF SUGAR BY AN AMMONIATED CUPRIC TEST GIVING REDUCTION WITHOUT PRECIPITATION.*

By F. W. PAVY, M.D., F.R.S.

To be able to effect the quantitative determination of a body with accuracy and facility is an important matter looked at in relation to the study of its bearings. In the case of sugar there are no reliable means of precipitating and weighing it, either alone or in combination, and thus in the chemical estimation of this principle an indirect method has to be resorted to. The only property upon which dependence can be placed, for the purpose of chemical quantitative analysis, is its reducing action, under the influence of heat, upon certain metallic oxides, and that of copper is the one which general experience shows to answer best.

In the ordinary volumetric application of the copper

* Abstract of a Paper read before the Royal Society Feb. 6, 1878.

* A Paper read at the Royal Society, January 16, 1879

test the precipitation and diffusion of the reduced suboxide through the liquid interferes with the clear perception of the precise point of complete decoloration, and thus detracts from its delicacy. For purposes where minute accuracy is of no moment, a sufficiently approximate result can be obtained, but for physiological investigation, and in other cases where precision is indispensable, the process is quite unfit for employment.

With the view of obtaining increased accuracy, chemists have had recourse to the plan of collecting the precipitate of reduced suboxide and weighing it as such or after reversion into the oxide. From the difficulty, however, that exists in procuring the metallic oxide in a pure and uniform state, and from the impossibility of completely freeing the filter paper used from adhering surplus copper solution, some uncertainty is given to the results obtained by this method. To obviate the difficulty here presented I suggested, in a communication published in the *Proc. Roy. Soc.* for June, 1877, that the precipitated suboxide should be collected and dissolved, and the copper frequently thrown down by the agency of galvanic action upon a platinum cylinder, as is now frequently done in the assaying of copper ores. The process has been found, as shown by the closeness observable in the results of counterpart analyses, to admit of the greatest precision, and I have turned it to extensive account in some recent physiological investigations I have conducted. In its application to such a purpose it may be held that time and labour should be considered as of no moment, but it frequently happens that a more ready process of investigation is needed than the gravimetric supplies, and on this account a volumetric method, free from the objection I have pointed out as belonging to the ordinary plan, constitutes a desideratum.

A few years back Bernard introduced, for physiological purposes, a modification of the ordinary volumetric process, which is attended with reduction and the non-precipitation of the reduced oxide. The process involves the employment of a large quantity of caustic potash, and the presence in the product to be tested of extraneous organic matter. Under these circumstances it happens that the reduced suboxide is held in solution instead of being allowed to fall, and thus decoloration without precipitation occurs and enables the point of disappearance of the colour of the test to be ascertained with precision. Bernard, in his remarks upon the test, simply made mention of the fact that under these conditions reduction without precipitation took place, but Dr. d'Arsonval,† his *Préparateur* at the College of France, refers the effect to the solvent influence of the extraneous organic matter in presence of the alkali.

Whilst engaged upon an enquiry into the merits of this test, the conclusion suggested itself to me that the agency preventing the deposition of the suboxide was the development of ammonia. With an absolutely pure solution of sugar, such as may be obtained by inverting the ordinary crystallised cane sugar (refined loaf sugar) no amount of potash will hinder the instantaneous precipitation of the suboxide. With commercial grape sugar, however, and in a still more marked manner with honey, interference with precipitation is exerted, and this, I am led to conclude, is due to the action of the potash in producing ammonia from the small quantity of nitrogenous organic matter incidentally present.

With this before me, the idea presented itself of resorting to the direct employment of ammonia for attaining the same result. It is well known to chemists that ammonia is a powerful solvent of the suboxide of copper, leading to the production of a perfectly colourless liquid; and this, from the facility with which it absorbs oxygen, quickly assumes a blue colour under exposure to air from the re-conversion of the cuprous into the cupric oxide.

If ammonia be added to the ordinary Fehling's solution, a liquid is obtained which is rendered colourless by boiling

with a sufficiency of sugar to effect the complete reduction of the cupric oxide present to the state of suboxide. As the saccharine product is dropped in the blue colour gradually fades, without any occurrence of precipitation to interfere with the perception of the precise moment when the point of complete decoloration is attained. The ammonia exerts no interference with the process of reduction, but simply dissolves the reduced oxide, leading, when complete decoloration is effected, to the production of a perfectly colourless, limpid liquid.

Enough ammonia must be present to secure that the suboxide is held in solution, and precaution must be taken that whilst the analysis is being performed the reduced oxide does not become re-converted into the oxide by exposure to the air. To obviate this the operation should be conducted in a flask instead of an open capsule.

The appliance that naturally suggests itself as most suitable for employment is a flask of about 80 cub. centims. capacity, with a cork inserted into the neck, through which a delivery tube from a Mohr's burette, graduated in tenths of a cub. centim., passes for dropping in the product to be examined. Through the cork, also, there must be an exit tube for the escape of air and steam from the flask. Should it be desired to avoid the impregnation of the surrounding atmosphere with ammonia, the exit tube may be connected by vulcanised tubing with a U-shaped tube containing fragments of pumice stone moistened with water or a weak acid. The burette being fixed in the stand, the flask is allowed to hang suspended, so that there may be nothing to obstruct the full view of the contents. The heat is applied by means of the flame of a spirit-lamp, and the best position for watching the disappearance of colour is by the light reflected from a white background specially provided for the purpose. It is convenient to have another burette, graduated in cub. centims., and of 100 cub. centims. capacity, fixed in the stand for holding and delivering the ammoniated copper solution. Messrs. Griffin, of Garrick Street, have constructed an arrangement to meet the requirements.

I at first took it for granted that in the action occurring the same relation existed between the amount of oxide of copper reduced and that of sugar oxidised, as under the employment of the copper test in the ordinary way, viz., that 5 atoms of oxide of copper were reduced by 1 atom of sugar, and the liquid I first employed was prepared by adding to 100 cub. centims. of Fehling's solution 300 cub. centims. of strong solutions of ammonia (sp. gr. 0.880) and 600 cub. centims. of distilled water. The liquid thus made contained one-tenth of Fehling's solution, and if it compared itself in the same manner as the latter, 10 cub. centims. of it would stand equivalent to 0.005 grm. of grape sugar. In working with this liquid the results obtained were so accordant in relation to each other that I had no misgiving about its uniformity of action; but I felt that before being definitely accepted they ought to be checked against known amounts of sugar. The accomplishment of this proceeding, however, is not altogether unattended with difficulty, on account of the uncertainty of obtaining grape sugar free from impurity and in a perfectly dried state.

The method I have adopted has been to operate upon weighed amounts of cane sugar and produce inversion by boiling with an acid. I first found that the cane sugar, which is sold in coarse colourless crystals—that which is known as "white crystal," and used for sweetening coffee—stood the test on examination for purity with Laurent's polarimeter. A weighed quantity was taken, and, after being inverted by boiling with hydrochloric acid, the acid neutralised, and the liquid brought to a known volume, subjected to treatment with the ammoniated copper liquid. Repeated trials were made with varying quantities, and it was found that the results stood in harmonious relation to each other, but that the amount of sugar indicated was larger than the calculated amount of invert sugar from the weighed quantity of cane sugar taken. At first I was at a loss for an explanation of this result, but subsequent

† *Gazette Hebdomadaire de Médecine et de Chirurgie*, September 14, 1877, p. 454.

observation has revealed that in the case of the ammoniated liquid, 6 atoms of oxide of copper are appropriated by 1 atom of sugar, instead of 5, as in that of Fehling's solution used in the ordinary way. When the reckoning is made upon this basis the results exactly correspond with the actual amount of sugar known to be present. Moreover, with solutions of ordinary grape sugar and diabetic sugar, examined comparatively with Fehling's solution used in the ordinary way and the ammoniated copper liquid, the results exactly accord under the reckoning that 5 atoms of oxide of copper are appropriated in the one case and 6 atoms in the other by 1 atom of sugar.

To be quite satisfied upon this point, a large number of observations under varying conditions have been made, and whilst what I have stated holds good for the ammoniated copper liquid prepared from Fehling's solution, without any further addition of alkali, and with the addition of potash to the extent of 1 grm. to 20 cub. centims. of the ammoniated test, yet a larger quantity of potash alters the action, and with 5 grms., and anything beyond, the behaviour is brought to the same as that of Fehling's solution used in the ordinary way, viz., 5 atoms only of oxide of copper are appropriated by 1 atom of sugar. With quantities of potash between the 1 and 5 grms., the results stand between the 5 and 6 atoms of cupric oxide.

I may mention that observation has further shown that whilst glucose prepared from starch behaves like other varieties of grape sugar, there is an intermediate product formed before the completion of the process of conversion, which behaves in a different manner from invert sugar, grape sugar, and sugar of diabetes. Estimations made with the ammoniated copper liquid coincide with those made with Fehling's solution without the presence of ammonia, and the addition of potash to the ammoniated liquid produces no modification of the result.

In order that the ammoniated copper liquid may be brought to the same standard of sugar value as Fehling's solution, and it is desirable that this should be the case, the proportion of copper must be increased so as to give 6 atoms against 5. By taking 120 cub. centims. of Fehling's solution, 300 cub. centims. of strong ammonia (sp. gr. 0.880) and making up to a litre with distilled water, the proper proportion is obtained, and the ammoniated liquid gives results corroborated in accuracy by the balance, and coinciding with those obtained by Fehling's solution employed in the ordinary way.

As a minor point it may be remarked that the diluted state presented by the ammoniated liquid offers an advantage by diminishing the liability to error arising from any want of absolute precision in measurement.

Twenty cub. centims. of the ammoniated copper solution, corresponding with 0.020 grm. sugar, having been run in from the burette containing the test, the flask is adapted to the cork attached to the delivery tube of the other burette containing the saccharine product for examination. The flame of a spirit-lamp is then applied underneath, and the contents of the flask brought to a state of ebullition and allowed to boil for a few minutes in order to get rid of the presence of air. The saccharine product is now allowed to drop from the burette until the blue colour of the test is just removed, and a perfectly colourless limpid state produced.

On account of the ammoniated copper solution used being only equivalent to 2 cub. centims. of Fehling's solution, it is necessary that the product to be examined should not be in too concentrated a form. For delicate observation it is convenient that the dilution should be such as to require the employment of from about 10 to 20 cub. centims. to decolourise the 20 cub. centims. of the ammoniated copper solution.

The ammoniated copper solution enjoys the advantage of possessing a self-preservative power. It is well known in the case of Fehling's solution that, in the course of time, not only does the liquid become impaired in stability, but actually reduced in strength, by the spontaneous

deposition of a certain amount of suboxide. Not so, however, with the ammoniated liquid. Here the conditions are such that under exposure to air the copper cannot fail to remain in solution and to be maintained in a fully oxidised state. A further advantage is given by the influence of the presence of ammonia on the colour of the test, for, in proportion to the height of colour of a volumetric liquid, so is its degree of delicacy as a reagent, and the effect of the addition of ammonia to the ordinary copper test is to considerably increase the blue colour belonging to it.

Seeing that the test here proposed acts with equal efficiency either in the presence or absence of extraneous organic matter, it is alike adapted for employment by the chemist, the physiologist, and the medical practitioner in relation to diabetes.

THE PREPARATION OF SINGLE REGULAR CRYSTALS OF ANY DESIRED SIZE.*

MR. FERDINAND MEYER, who has for thirty years studied the conditions necessary for obtaining large, single, and regular crystals of chemical salts, has published his method in the *Archiv der Pharmacie*, October, 1878, from which we take the following:—

Prepare a solution of any salt in water of such a strength that, after standing twenty-four hours, a portion of the salt will separate in crystals. Pour off the mother-water, select a few of the best formed crystals, and place them on a plate of glass, which lies in a rather tall vessel. Then re-dissolve a little of the dry salt in a small quantity of the mother-water, add this supersaturated solution to the main bulk of the mother-water, pour this upon the crystals on the plate of the glass, and place the vessel into a room where the temperature remains as uniform as possible, best in the cellar. The temperature of the room should be ascertained by a thermometer, and in case of any changes of temperature, a further quantity of the salt is to be dissolved in the mother-water. This must be repeated every twelve or fourteen hours, until the crystals have reached the desired size. If the solution is too strong, single regular crystals are seldom obtained at once, but this is generally of no consequence, for, as long as one side at least is perfectly formed, it is only necessary to turn them two or three times to cause the other sides likewise to become perfect. As the crystals increase in size care must be taken to give them a correct position in the plate of glass: and, if the solution is at all concentrated, the crystals must be carefully freed from adhering irregularities, and then replaced in the solution.

In a solution of alum, a very oblique octoëder is usually obtained first. This may be allowed to reach a considerable size, after which it is to be laid successively on the narrow sides, when it will gradually become a regular octoëder. If it is always kept lying on the broadest sides it will continue to grow obliquely.

It is well known that several isomorphous salts may be crystallised, one over the other, in layers, without a change of crystalline form. Chrome-alum crystals may thus be covered with crystals of ordinary alum. The largest crystal of this kind obtained by the author weighed over three pounds.

The author also observed that, when employing the same mother-water for a considerable time, the crystals began to show blunt or flattened points. This happened with regular as well as with oblique crystals, so that in place of eight surfaces, the regular crystals gradually assumed sixteen equal sides, and the irregular ones, fourteen smaller and two larger sides. If, however, the mother-water be acidulated with a little sulphuric acid, this flattening of the point occurs but rarely.

* From "New Remedies," January, 1879.

As a general rule, this changing the position of the single crystals of any salt, but particularly of sulphate of zinc, copper, nickel, or magnesium, and of Rochelle salts, different forms of the same system are obtained. If crystals of Rochelle salt, which may easily be obtained of large size, are always placed upon one and the same side, one-half of the crystal will become perfectly developed; but if they are laid, alternately, upon the two opposite long surfaces, the development is less regular. On placing large crystals of the same salt, even if only half developed, lengthwise into the liquid, alternately upon either end, development of the lateral surfaces proceeds very regularly.

It is not advisable to introduce crystals into a solution if the latter is at all warm, or to pour a warm solution into a cold one containing crystals, as the latter are thereby generally torn or broken. If a portion of a crystal has been by accident broken off, it may be repaired by subjecting it to the above-detailed process. In a crystal of chrome alum, from which a piece weighing ten grms. had been broken off, the gap was completely restored by subjecting it to the feeding process for a fortnight.

As soon as the crystals have attained the desired size it is best to place them into less concentrated solutions, in a slightly cooler place; this after-treatment causes the surfaces to become smooth and levelled, and the edges to become sharp.—*Pharmaceutical Journal*.

ANALYSIS OF BLEACHING-POWDER.

By GEORGE WHEWELL, F.I.C., F.C.S.

BEING requested some time ago to make an analysis of an unsatisfactory sample of commercial bleaching-powder, and finding it to absorb moisture very rapidly, I made the following experiments, which I think might interest your readers.

The rate of absorption was determined by exposing 100 grs. to the atmosphere of the laboratory and weighing every two hours during the day.

In 24 hours it absorbed 25 per cent of moisture.

" 48	"	"	41	"	"
" 72	"	"	52.3	"	"
" 96	"	"	64.9	"	"
" 103	"	"	69	"	"

A small quantity now left is in a liquid condition.

The sample was sold as containing 35 per cent of available chlorine, but was found to only contain 28.6 per cent. Its percentage composition was:—

Calcium..	..	..	..	..	30.88
Chlorine	..	..	..	..	58.13
Difference (oxygen, &c.)	..	..	..	..	10.99

100.00

The calcium was determined as carbonate, and the total chlorine as chloride of silver.

ALUM IN FLOUR AND BREAD.

By M. D. PENNEY, F.C.S.

THE quantity of alumina naturally present in pure flour and bread has been a subject of discussion for many years, and has been erroneously assumed by some chemists not to exceed an amount equal to 8 or 10 grains of alum per 4 lbs. At a meeting of the Society of Public Analysts, February 5, 1875, it was stated by one speaker to be as low as 2 grs., and by others as high as 10 grs., expressed as alum per 4 lbs. of bread; and a rather prominent analyst asserted at the "Selby case" a few months ago that, in his opinion, 2 grs. were quite sufficient to allow for. It has recently

become apparent that the attempt to fix upon a standard must be abandoned, and perhaps some experiments made by me last April and May, may assist to throw more light on the question.

TABLE I.
Six Samples of Flour.

No.	Grains per 4 lbs.		
	Alumina expressed as Alum.	Ferric Oxide.	Siliceous Matter.
1.	24.30	5.47	50.00
2.	21.75	6.06	55.00
3.	21.25	4.90	56.00
4.	17.00	5.20	40.00
5.	12.40	5.30	30.00
6.	6.34	4.77	22.00

In the first table are the analyses of six pure flours. I was satisfied that they were quite free from alum, and yet I found alumina equal in No. 1 equal to 24.30 grs. of alum per 4 lbs.

The first four flours came from a large mill where much foreign grain is used, and No. 6 was made at a small country mill from English grown wheat alone. It occurred to me that the quantity of alumina in flour might depend on the variety of grain from which it was prepared; and with a view to ascertain if such were the case, I made several analyses of wheats from different countries, and have tabulated the results.

TABLE II.

No.	Variety of Wheat.	Milligrams. per 100 grms.		
		Phos. of Alumina.	Phos. of Iron.	Siliceous Matter.
1.	Calcutta	24.30	18.10	164.00
2.	Ditto	21.00	15.16	184.00
3.	Ditto	18.50	20.00	170.00
4.	Kourish	27.20	34.00	206.00
5.	Ditto	31.00	36.00	226.00
6.	Russian	17.20	20.00	101.00
7.	Ditto	24.45	17.20	131.00
8.	Ditto	13.10	11.40	70.00
9.	Ditto	16.35	16.34	126.00
10.	Chicago	4.00	9.00	48.00
11.	Oregon	4.00	8.11	36.00
12.	English	5.12	7.40	35.00
13.	Ditto	6.40	14.20	40.00
14.	Ditto	7.30	14.00	30.00
15.	Ditto	3.80	4.70	32.00
16.	Stetin	16.33	26.40	109.00
17.	Ditto	13.24	21.23	120.00
18.	California	3.00	2.21	31.00
19.	11 varieties mixed	15.10	12.30	59.00
20.	Egyptian as impd.	167.00	64.50	370.00
21.	Ditto, hand picked	49.49	27.20	117.00
22.	Ditto, washed ..	14.10	6.36	101.00

Nos. 1, 2, 3 are Calcutta wheats, now largely imported and much prized by millers. The following six refer to Russian wheat. The Kourish variety is particularly high in alumina and silica. Nos. 10 and 11, and the English wheats, Nos. 12 to 15, are very much alike. No. 18 was a sample of Californian wheat, a pretty looking, large, white grain, remarkably low in alumina and iron oxide. No. 19 was a mixture of 11 varieties mixed by a miller preparatory to grinding. Egyptian wheat, as imported, is much intermixed with clay. It seems that the lighters in which it is carried on the Nile are unprovided with hatches, and in lieu of them the cargo is covered with mud, which is soon baked hard in the sun and gives a firm footing to the boatmen. This rough treatment accounts, no doubt, for its presence in the form of dust, pellets, and lumps. It is always more or less cleansed preparatory to grinding; but it does not appear practicable, with the cleansing apparatus attached to the best constructed mills, to get rid of all the extraneous matter in

grain. Sometimes the holes made by the weevil (an insect infesting some kinds of wheat, Calcutta for instance) are filled with dirt, and no amount of working or screening will extract it.

A glance at the table will show what kind of flour to expect from any of the wheats. It is rarely, however, that any one kind is ground by itself. It is the practice of millers to mix several kinds, and the mixture varies in every mill and in every part of the county according to the supply of foreign grain and the quality of our home-grown.

No. II. Table shows that some varieties of wheat are more contaminated than others with alumina. I use the word contaminated advisedly, as I believe that the greater portion of it is due to extraneous matter, and it reveals the source of the excessive quantity of alumina occasionally met with in flour.

Since it is impossible to fix upon a standard of natural alumina, and since it is erroneous to regard its amount, even in the presence of alum, as in any degree a measure of alum, it may be asked—What is the use of estimating it at all?

I venture to suggest that it might indicate whether the grain was or was not cleansed properly before grinding; but the estimation of silica would answer just as well. Experience will enable us, perhaps, to decide on the limit to allow for clay in flour. If a line were drawn somewhere it would probably make millers more careful in the cleansing process.

Dr. Dupré's "chloroform" method of separating alum from flour appears from his reports to be satisfactory. A correct quantitative test for alum in bread is still a desideratum. As for a qualitative test, there cannot be anything more simple and certain than the logwood test properly applied. I have tried it repeatedly with different kinds of flour and bread, and never found it to fail. It does not matter, in my opinion, whether the solution is new or old, and all the logwood chips I have come across answer well. I have never found any difficulty in distinguishing the alum colour from that caused by soil, magnesia, or any probable constituents of flour or bread. I therefore place implicit trust in it, and feel rather surprised that its efficiency is sometimes questioned.

Hull, January 31, 1879.

ON THE FORMATION AND COMPOSITION OF SOME COMPLEX OXIDES OF COBALT AND NICKEL.*

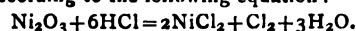
By THOMAS BAYLEY, Assoc. Royal Coll. of Science, Ireland.
Demonstrator of Practical Chemistry, Analysis, and Assaying in the
Mining School, Bristol.

WHILE preparing standard solutions of nickel and cobalt salts for the purposes of a research on the colorimetric relations and on the colorimetric estimation of those metals, I was endeavouring to use a modification of the method of estimating nickel and cobalt, indicated by Bunsen†, depending for their determination on the iodine liberated by the higher oxides of these metals in contact with hydrochloric acid and potassium iodide. The method was as follows:—The solution of the nickel or cobalt salts was made alkaline by soda and then mixed with excess of sodic hypochlorite obtain by the action of cold dilute sodic carbonate on fresh bleaching-powder.

After allowing the slightly warm solution of nickel or cobalt to stand some time, so as to ensure complete oxidation, the temperature was raised until brisk effervescence ensued, and the solution allowed to remain at that temperature until the excess of hypochlorite was decomposed. When the evolution of oxygen had ceased, the liquid was

boiled for about half an hour. I found that by this process, it is easy to destroy all matter, except the oxide, capable of liberating iodine on treatment with potassic iodide and hydrochloric acid. The solution having been cooled, it was mixed with excess of potassic iodide, and then with enough hydrochloric acid to dissolve the suspended oxide. The liberated iodine was then estimated by a standard solution of sodic thiosulphate ($\text{Na}_2\text{S}_2\text{O}_3$).

In the first experiments I used a standard solution of nitrate of nickel, and calculated the nickel from the iodine set free according to the following equation:—



The results were not satisfactory, as will be seen from the following Table:—

Nickel used.	Nickel found.
Grm.	Grm.
0.1570	0.1437
0.1570	0.1580
0.1570	0.1465
0.1570	0.1568
0.1570	0.1639
0.1570	0.1541

Besides these analyses there were several which yielded a far less quantity of nickel. The same method was then applied to cobalt, with this difference, that the solution was boiled only for a few minutes, as I found that length of time sufficient for the decomposition of the last traces of hypochlorite. The amounts of iodine liberated were much greater than would be due to the oxide Co_2O_3 , while they agreed perfectly with an oxide Co_3O_5 , thus:—

Cobalt used.	Iodine liberated.	Theory of Iodine for Co_3O_5 .
Grm.	Grm.	Grm.
0.1865	0.5338	0.5343
0.1865	0.5380	0.5343
0.1865	0.5328	0.5343

I now repeated the experiments with nickel, taking care to boil the liquid only a minute or two. In one or two instances it was not boiled, but the precipitate filtered off and washed. The results were as follows:—

Nickel used.	Iodine liberated.	Theory of Iodine for Ni_2O_3 .
Grm.	Grm.	Grm.
0.1570	0.4428	0.4521
0.0785	0.2263	0.2260
0.1835	0.5318	0.5284
0.1570	0.4490	0.4521

With a mixture of 0.1863 gram. cobalt, and 0.1835 gram. nickel:—

Iodine found.	Theory for Iodine due to Ni_2O_3 & Co_3O_5 .
Grms.	Grms.
1.0532	1.0627

In the last cases the oxides were not boiled, but the solution was allowed to stand over the steam bath for a few hours.

With solutions of known quantities of nickel I now made the following experiments. The solution with the suspended oxide was boiled for some hours.

Nickel used.	Iodine found.	Theory of Iodine for Ni_2O_3 .
Grm.	Grm.	Grm.
0.1835	0.2009	0.2642
0.1835	0.1812	0.2642
0.1835	0.2321	0.2642
0.1835	0.3310	0.2642
0.1835	0.2830	0.2642
0.1835	0.3765	0.2642
0.1835	0.1069	0.2642

In the last experiment the solution was boiled for a few days.

* A Paper read before the Royal Irish Academy, June 25, 1877.
† *Ann. Ch. Pharm.*, lxxvi., 265.

A quantity of cobalt nitrate was now mixed with soda and sodic hypochlorite, and allowed to stand in a warm place until effervescence had ceased. The precipitated oxide of cobalt was then well washed with warm water and dried, till constant, under the air pump, over strong sulphuric acid.

A sample of this oxide was then submitted in succession to various temperatures. The results were as follows:—

Grm.	Grm.
Oxide taken 0.9278 ($\text{Co}_3\text{O}_5, 4\text{H}_2\text{O}$)	
Oxide dried at 100°C . } 0.8760 Theory for $\text{Co}_3\text{O}_5, 3\text{H}_2\text{O}$..	0.8770
Do., 138°C . } 0.8223 " " $\text{Co}_3\text{O}_5, 2\text{H}_2\text{O}$..	0.8263
Do., 310°C . } 0.7548 " " $\text{Co}_3\text{O}_5, \text{H}_2\text{O}$..	0.7756

After this experiment there was an appearance of change on the surface of the oxide.

Grm.	Grm.
Oxide after ignition to redness ..	0.6840 Theory for Co_3O_4 ..
	0.6798

Another portion of the same sample:—

Grm.	Grm.
Oxide dried at 100°C . 0.4070 ($=\text{Co}_3\text{O}_5, 3\text{H}_2\text{O}$).	
Oxide after ignition to redness ..	0.3125 Theory for Co_3O_4 ..
	0.3154

Another sample prepared in the same way, but left longer over the air pump:—

Oxide taken dried over H_2SO_4 .. 0.7083.

The oxide was now heated to low redness in a tube in a current of dry oxygen, and the water given off collected in a tube filled with calcium chloride. The oxide was afterwards ignited to bright redness in air.

	Grms.
Calcium chloride tube ..	65.3250
Calc. chl. tube + water ..	65.4680
	0.1430 = OH_2

	Grm.
Oxide after ignition in a tube ..	0.5493
Oxide after ignition in air ..	0.5293

The results of this experiment are compared below with the theory for $\text{Co}_3\text{O}_5, 4\text{H}_2\text{O}$. For a reason which will be seen further on, I have added the theory for $\text{Co}_2\text{O}_3, 3\text{H}_2\text{O}$.

Theory for $\text{Co}_3\text{O}_5, 4\text{H}_2\text{O}$.	Found.	Error.
Grm.	Grm.	
$4\text{H}_2\text{O}$..	0.1549	0.1428 .. -0.012
Co_3O_5 ..	0.5534	0.5493 .. -0.004
Co_3O_4 ..	0.5190	0.5293 .. +0.013

$\text{Co}_3\text{O}_5, 4\text{H}_2\text{O}$.	Found.	$\text{Co}_2\text{O}_3, 3\text{H}_2\text{O}$.
Grm.	Grm.	Grm.
OH_2 ..	21.86	24.54
Co_3O_4 ..	73.27	73.05
	95.13	97.59

Winkelbleck obtained an oxide in the same way as I prepared my samples, only that he boiled his with strong potash before washing it. He dried his oxide over strong sulphuric acid. According to him, the formula is $\text{Co}_2\text{O}_3, 3\text{H}_2\text{O}$. His results were as follows:—

	Per Cent.	Per cent.
	(I.)	(II.)
2Co ..	53.83	53.93
3O ..	21.62	21.46
$3\text{H}_2\text{O}$..	24.26	24.61
		53.64
		21.82
		24.54

The question, which is the true formula of the oxide I obtained, is determined by the amount of iodine liberated by the oxide on treatment with potassic iodide and hydrochloric acid. According to the formula $\text{Co}_2\text{O}_3, 3\text{H}_2\text{O}$, there should be liberated 0.402 grm. by 0.1865 grm. of cobalt; according to the formula $\text{Co}_3\text{O}_5, 4\text{H}_2\text{O}$, 0.5343 grm. should be liberated. I found in three experiments 0.5338, 0.5380, and 0.5328.

When the oxide $\text{Co}_3\text{O}_5, 4\text{H}_2\text{O}$, obtained as described above, is boiled for an hour or two in the solution in which it is precipitated, and the amount of iodine liberated then estimated, the result points to the formation of the oxide $\text{Co}_{12}\text{O}_{19}$, intermediate between Co_3O_5 and Co_2O_3 .

Co Taken.	Theory of I for Co_3O_5 .	Theory of I for Co_2O_3 .	Found.
Grm.	Grm.	Grm.	Grm.
0.1865	0.5343	0.4007	(1) 0.453
	Mean.		(2) 0.463
	0.4676		(3) 0.462
	Theory for $\text{Co}_{12}\text{O}_{19}$.		Found.
	Iod.		Grm.
0.0620	0.1566 grm.		0.1556

In the last experiment a fresh solution of cobalt, and a fresh solution of potassium bichromate (to standardise the thiosulphate), were used.

A quantity of the oxide of cobalt prepared by precipitating with potash and sodic hypochlorite, and boiling for some hours, then washing and drying over sulphuric acid *in vacuo*, was submitted to a current of air at a low red heat, and the water collected and weighed in a calcium chloride tube. The oxide was afterwards ignited to bright redness in air.

	Grms.
Oxide dried over H_2SO_4 ..	0.7455
Oxide after ignition in tube ($\text{Co}_{12}\text{O}_{19}$) ..	0.6255
Oxide after ignition in air (Co_3O_4) ..	0.5975
Calcium chloride tube + OH_2 ..	65.5930
Calcium chloride tube ..	65.4645
	0.1255 = OH_2

	Grm.	Found.
Theory for $\text{Co}_{12}\text{O}_{19}, 11\text{H}_2\text{O}$.		Grm.
OH_2 ..	0.1218	0.1255
$\text{Co}_{12}\text{O}_{19}$..	0.6236	0.6255
Co_3O_4 ..	0.5940	0.5975

Percentages.

	Theory for $\text{Co}_{12}\text{O}_{19}, 11\text{H}_2\text{O}$.	Found.
	Grm.	Grm.
OH_2 ..	16.34	16.83
$\text{Co}_{12}\text{O}_{19}$..	83.65	83.90

On attempting to prepare Ni_3O_5 in the dry state by precipitating, washing, and drying *in vacuo*, I found that the moist precipitate gave off oxygen as soon as the liquid in which it was precipitated was removed. The moist precipitate was allowed to stand some days, and then left over the air-pump for about a week, in order to allow time for this change to be complete. Owing to some interruption, I have as yet had time to prepare only one sample by this means. The results of the analysis agree closely with the formula $\text{Ni}_8\text{O}_{12}, 9\text{H}_2\text{O}$, one-ninth of the water being lost at 100°C .

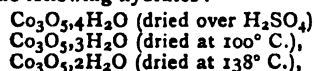
	Oxide taken.	Theory of Iodine for $\text{Ni}_8\text{O}_{12}, 9\text{H}_2\text{O}$.	Found.
	Grm.	Grm.	Grm.
Dried over H_2SO_4 ..	0.1705	0.1607	0.1661
" " " ..	0.2012	0.1896	0.1895
" " " ..	0.2375	0.2238	0.2243
Dried at 100°C . ..	0.2080	0.2005	0.2026

The water in this oxide was determined by igniting the oxide in a platinum boat in a combustion-tube, and weighing the water lost by means of a calcium chloride tube.

Oxide taken.	Theory for $\text{Ni}_8\text{O}_{12}, 9\text{H}_2\text{O}$.	Found.
Grms.	Grms.	Grms.
0.8723	0.1748 grm. OH_2	0.1775
	Per Cent.	
OH_2	20.039	20.34
	Per Cent.	
	Theory for $\text{Ni}_8\text{O}_{12}, 9\text{H}_2\text{O}$.	Found.
	Grms.	Grms.
NiO	74.02	(1) 74.15
		(2) 74.29

Summary.

Under the influence of the hypochlorite solution nickel and cobalt form the oxides Ni_3O_5 and Co_3O_5 . On boiling the liquid containing Co_3O_5 it loses oxygen, and passes to the form $\text{Co}_{12}\text{O}_{19}$, intermediate between Co_2O_3 and Co_3O_5 . Under similar circumstances, Ni_3O_5 appears to decompose without forming stable lower oxides, although it is probable from the results that the formation of Ni_2O_3 is a stage in the process. The oxides Co_3O_5 and $\text{Co}_{12}\text{O}_{19}$ appear to be stable at a low red heat: they are distinguished from Co_3O_4 by a slight difference of colour. Co_3O_5 has the following hydrates:—

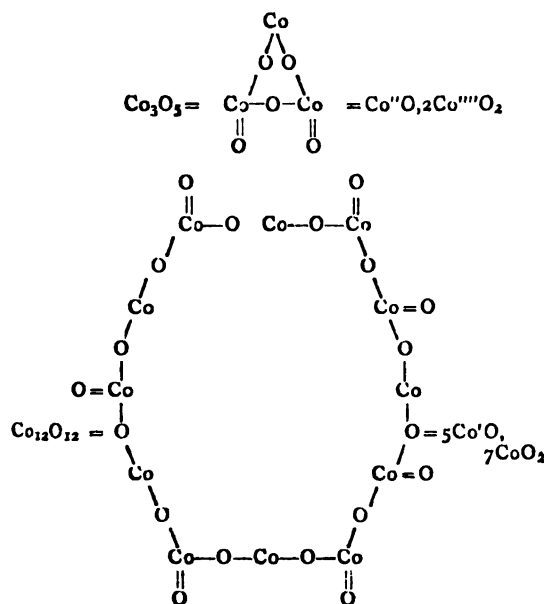


and probably,



Ni_3O_5 decomposes while still moist when its precipitating liquid is removed by washing. In the one experiment which was made, the resulting dried compound agreed closely with the formula $\text{Ni}_8\text{O}_{11} \cdot 9\text{H}_2\text{O}$. I have found that when $\text{Co}_3\text{O}_5 \cdot 4\text{H}_2\text{O}$ is treated with cold dilute nitric acid part is dissolved with evolution of oxygen, and that part remains insoluble. I hope, in a future paper, to give the results of some similar experiments undertaken for the purpose of determining the proximate constitution of these oxides.

It may at first sight appear that the formula $\text{Co}_{12}\text{O}_{19}$ is inadmissible on account of its complexity, but as the iodine method clearly shows that the oxide is exactly intermediate between Co_3O_5 and Co_2O_3 , and as the formula $\text{Co}_{12}\text{O}_{19}$ is the simplest formula for such an oxide, it would seem that we must accept it, especially when we consider the tendency of cobalt to form compounds vieing in complexity with many of the products of organic chemistry. It may be that the application of the iodine method to the examination of the oxides of other metals would lead to the acceptance of formulæ more complex than those now admitted. The two oxides of cobalt described in this paper, and indeed other oxides of cobalt, may be represented graphically by rings somewhat analogous to the well-known benzene "ring" of the aromatic carbon compounds and to the zinc "ring." Thus—



This investigation was conducted in the Chemical Laboratory of the Royal College of Science, Dublin.

Further Note.—I have found by experiment that when any of these peroxides are decomposed by hydrochloric acid, and the evolved chlorine passed into a solution of potassic iodide, the amount of iodine liberated is exactly equal to that which is liberated by decomposing the same weight of the peroxide in a solution containing potassic iodide. The presence of caustic alkali in large quantity appears to retard the passage of Co_3O_5 into $\text{Co}_{12}\text{O}_{19}$ during the boiling process. Co_3O_5 , indeed, seems to be more stable in proportion as the amount of free alkali is greater.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 4, January 27, 1879.

Third Reply to M. Berthelot.—M. Pasteur.—A continuation of the wearisome Claude Bernard controversy.

Researches on the Relations of Spectral Analysis with the Solar Spectrum.—J. N. Lockyer.—The substance of this paper will be found in the *CHEMICAL NEWS*, vol. xxxix., pp. 1, 11.

Composition of the Banana and the Utilisation of this Fruit.—V. Marcano and A. Muntz.—In addition to the production of its fruit the banana tree is serviceable for keeping the surrounding soil moist in dry seasons, a matter of great importance in the cultivation of coffee. The fruit when ripe contains 10 per cent of cane-sugar and 3 per cent of inverted sugar.

Application of His Atomic Theory to Certain Minerals.—A. Gaudin.—Nothing short of a full translation could give a satisfactory view of the speculation contained in this paper.

Displacement of Spectral Rays Due to the Rotatory Movement of the Sun.—L. Thollon.—The author by making use of his newly invented spectroscope has observed a displacement of the solar rays perfectly well defined, and evidently approaching to the results of calculation.

Radiation of Incandescent Platinum.—J. Violle.—The author has made a number of measurements of the intensity of the red light emitted by platinum at different temperatures, from 900° to 1775° , the melting-point of the metal. The intensity of the red light, scarcely sensible at 500° , increases at first very rapidly and then more slowly. At 2910° its intensity is not greater than at the melting-point of silver.

Illumination of the Lines of Molecular Pressure and the Trajectory of Molecules.—W. Crookes, presented by Th. du Moncel.—The substance of this paper has appeared in the *CHEMICAL NEWS*, vol. xxxviii., p. 279.

Electro-dynamic Phenomena and in Particular on Induction.—H. de Meaux.—The author concludes that in a closed circuit the intensity of a given current cannot be changed by the induction of an indefinite cylindrical conductor upon another of the same form by surrounding one or other of these conductors, or even both, with a concentric metallic covering communicating with the ground in its whole length.

A New Bell's Telephone, with a Loud Voice.—M. Gower.—The peculiarities of this instrument are said to be due to the fact that the poles are placed as in Faraday's electro-magnet. The diaphragm is thicker, larger, and tighter than those commonly constructed. The case containing the whole is metallic, and forms a sounding-chest, and is furnished with a bell mouth which amplifies sound

On Amalgams of Chrome, Manganese, Iron, Cobalt, and Nickel, and a New Process for Preparing Metallic Chrome.—H. Moissan.—If a concentrated solution of chromous chloride is shaken up in water with sodium amalgam, a part of the sodium decomposes water, gives off hydrogen, and forms soda, which precipitates a certain quantity of chrome. The other part of the amalgam produces by double decomposition chloride of sodium and an amalgam of chrome. To be certain that no sodium remains in the mercury the amalgam obtained is kept for an hour in boiling water, stirring from time to time. If this amalgam is heated in a current of hydrogen to 350° metallic chrome is left as a black amorphous slightly coherent substance. The same method serves for preparing amalgams of manganese, iron, cobalt, and nickel.

Preparation of Methyl-formic Ether and Pure Methylic Alcohol.—MM. Bardy and Bordet.—Not suitable for abstraction.

Active Principles of *Sarracenia Purpurea*.—F. Hétet.—From this plant the author has extracted an alkaloid whose characters agree with those of veratrin. It is curious that the *Sarracenia*, though belonging to a remote family, should contain a poison similar to that of colchicum.

Physiological Action of so-called "Garnet"—the Residue of the Manufacture of Magenta.—M. Jousset de Bellesme.—This substance dissolved in treacle is much more used for colouring wines than is pure magenta, which would give a shade too much bordering upon a rose. In a number of experiments made upon animals this colour always produced fatal results. One c.c. proved fatal to a cat in twelve days. The symptoms are persistent diarrhoea and rapid wasting.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin
No. 14, 1879.

Action of Hydrochloric Acid upon Certain Double Salts of Sulphuric Acid.—C. Hensgen.—The author divides the double sulphates into three classes: those in which each of the sulphates present is decomposable by hydrochloric acid; those in which one of the sulphates only is attacked; and those in which neither of the component sulphates is affected. He has, in the first place, studied the behaviour of potassio-cupric sulphate, of potassic iron, alum, and of ammonio-ferrous sulphate.

Iso-heptylic Acid from β -Hexyl-iodide.—Otto Hecht and J. Munier.—Iso-heptylic acid is a limpid oily liquid of faintly rancid odour, sparingly soluble in water, and boiling at 211° to 213°.

Synthesis of Meta-nitro-cinnamic Acid.—R. Schiff.—The author obtains this acid by heating a mixture of equal molecules of nitro-benzaldehyd, anhydrous acetic acid, and sodic acetate for eight hours in a vessel provided with an ascending cooler.

On Acetylen Urea.—C. Böttiger.—A further examination of bodies which the author obtained last year on treating urea with glyoxal in presence of hydrocyanic acid.

On Hydroxylation by Direct Oxidation.—R. Meyer.—Not susceptible of useful abstraction.

On Oxypropyl-benzoic Acid and its Derivatives.—R. Meyer and J. Rosicki.—A closer examination of the acid in question, which one of the authors recently obtained by the oxidation of cuminic acid.

Explanation on the Formula of Uric Acid.—R. Fittig.—A question of priority.

New Analyses of the Mineral Springs of Passug, Solis, and Tiefenkasten in the Canton Grisons, Switzerland.—A. von Planta-Reichenau.—Six pages of analytical results; possibly of medical interest.

Specific Rotation of Cane-sugar.—B. Tollens.—An important paper, but not susceptible of useful abstraction.

Further Observations on Pyrrol and its Derivatives.—Chichester A. Bell.—An examination of the so-called ethyl-pyrrol of Lubawin, and of its behaviour with hydrochloric acid and with bromine.

Camphen Derived from Camphor, and the Synthesis of its Homologues.—F. V. Spitzer.—This paper treats of the camphen obtained from the most infusible camphor-bichloride, and on ethyl-camphen and isobutyl-camphen.

Camphor Chlorides.—F. V. Spitzer.—The camphor-bichloride discovered by the author is readily soluble in alcohol and ether. It melts at 155° to 155.5°. If preserved in a somewhat moist condition it easily gives off hydrochloric acid.

Cinchonic Alkaloids.—A. Claus.—Not adapted for abstraction.

Imido-thio-ether.—A. Pinner and F. Klein.—This product is obtained by a reaction to that which yields imido-ether.

Reimann's Färber Zeitung,
No. 1, 1879.

This issue gives a very dismal account of the condition of the once so prosperous and celebrated print-works of Alsace.

No. 2, 1879.

A silk merchant in Zurich is somewhat roughly handled for having advertised his silks as "pure" and for having called public attention to the weighting system, which Dr. Reimann declares is carried "only to the extent of 150 per cent."

Since the cultivation of madder in France has dwindled to about one-eighth of its former extent it is proposed to abolish the red trousers of the French army.

There is the commencement of a review of the specimens of calico-printing at the late Exhibition. Several English exhibitors are praised, but we note the remark that none of the English furniture-prints display the fiery red which is so admired in French work.

No. 3, 1879.

Following up the example about to be set by France it is proposed that the blue coats and grey trousers of the German army should be abolished, as they necessitate the use of indigo, and that alizarin browns or blacks should be used in their stead.

No. 4, 1879.

The principal articles in this issue are a dissertation in favour of protection and complaints of injustice, or, at least, negligence, in the award of prizes at the late Paris Exhibition.

It is announced that the financial condition of the Berlin Dyers' Association is very unsatisfactory.

No. 5, 1879.

There is here an account of the sophistication of soaps with rosin, potato-flour, silicate of soda, and alumina, all of which additions except the last are declared to be positively injurious.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 3, January 16, 1879.

The Telemachon.—W. Wallace, of Connecticut, is said to have invented a new apparatus under this name by which motive power can be conveyed many hundred miles without loss or difficulty.

The Teleroscope.—M. Senlecq, of Ardres, has submitted to the examination of M. du Moncel a project for re-producing telegraphically at a distance images obtained

in the camera. The apparatus is based upon the property possessed by selenium of offering an electric resistance variable according to the degrees of light.

No. 4, January 23, 1879.

A man is said to have died in great torment from having struck a match upon his thumb-nail, when a minute fragment of phosphorus lodged under the nail and brought on a violent inflammation, which extended to the shoulder.

According to the *Revue Britannique* ether is in common use as a stimulant in Drapers' Town, in Ireland, where its odour can be recognised at the distance of half a mile! The grass in Hyde Park is said to be strewn with empty ether-bottles which ladies of fashion have thrown out of the windows of their carriages.

M. Audouin recommends chromic oxide as the most refractory material known for fire-bricks, crucibles, &c. It resists the highest known temperatures, and is attacked neither by oxides of iron nor by silica.

No. 5, January 30, 1879.

According to Dr. Draper the rain-fall at New York has been continuously decreasing since 1869.

No. 6, February 6, 1879.

French Production of Wine.—The total production of wines for 1878 was 48,000,000 hectolitres, and that of cyder and perry 12,000,000. Half a million hectares of vineyards have been annihilated by the phylloxera.

It appears that "green oysters" are sometimes fraudulently prepared by steeping them in a solution of a salt of copper.

Simple Method of Detecting Magenta in Wines, Fruit-syrups, &c.—Prof. Flückiger adds to portions of the sample, dilute if needful, recently-prepared chlorine and bromine water. If genuine the colour in both becomes a faint yellow. If magenta is present chlorine produces a deep dirty tint and bromine a violet colour, followed in time by the formation of violet flocks. Neither reagent destroys the colour.

La Lancette Belge, January 15, 1879.

According to statistical returns, sulphate of copper has caused the death of 77 persons in Belgium during the last ten years. The phosphorus obtained from matches has proved fatal to 170, verdigris to 33, sulphuric acid to 30, and cantharides to 24.

MEETINGS FOR THE WEEK.

- MONDAY, 24th.—Medical, 8.
— Royal Geographical, 8.30.
— Society of Arts, 8. "Dwelling Houses: Their Sanitary Construction and Arrangements," by Dr. W. H. Corfield, M.A. (Cantor Lectures.)
— London Institution, 5.
TUESDAY, 25th.—Civil Engineers, 8.
— Royal Institution, 3. "Animal Development," Prof. Schäfer.
— Anthropological, 8.
WEDNESDAY, 26th.—Society of Arts, 8. "Indian Pottery at the Paris Exhibition," Dr. Birdwood, C.S.I.
— Geological, 8.
THURSDAY, 27th.—Royal, 8.30.
— Royal Institution, 3. "Sound," Prof. Tyndall.
— Philosophical Club, 6.30.
— London Institution, 7.
FRIDAY, 28th.—Royal Institution, 9. "The Sorting Demon of Maxwell," Sir W. Thomson.
— Quekett, 8.
SATURDAY, 29th.—Royal Institution, 3. "Lessing," by Reginald W. Macan.

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THE CHEMICAL NEWS.

Vol. XXXIX. No. 1005.

ON THE LIQUEFACTION OF GASES.*

By J. J. COLEMAN, F.I.C., F.C.S.

THREE methods have been employed in the liquefaction of gases:—I. Increase of pressure at normal temperatures. II. Abstraction of heat at normal pressures. III. The combination of these two operations. The gases which can be liquefied by pressure at normal temperatures are limited. Natterer showed us, some years ago, that under such conditions the enormous pressure of 3000 atmospheres was insufficient to liquefy hydrogen and oxygen. It was reserved for our countryman, Dr. Andrews, to demonstrate the fallacy of relying upon pressure alone for gaseous liquefaction. To use the words of Prof. Wurtz in his recent Faraday Lecture, he has shown us that with all vapours there is a point at which the molecular movements caused by heat finally gain the victory over the force of cohesion, whatever be the pressure to which the air is subjected. Dr. Andrews describes it as the "critical point;" Mendelejeff names it the "absolute boiling-point."

It is, therefore, easy to understand why Natterer failed and Cailletet and Pictet succeeded with their experiments. Indeed, it would appear from these researches, and the principles underlying them, and if it be taken for granted that the molecules of all gases have a tendency to gravitate to each other independent of external pressure, that of the two agents for effecting the liquefaction of gases the abstraction of heat is more efficacious than the application of pressure.

This is evident, and follows from the ordinary laws of gaseous expansion. It is a well-established fact that gases expand uniformly, and that the rate of expansion is such that a given volume of gas at the freezing-point of water will become double its bulk at 491° F. Therefore, if it be followed in its path of contraction for temperatures below freezing-point, by the time its temperature becomes reduced 491°, which is the absolute zero of physicists, the gas must necessarily be either liquefied or solidified. The greatest artificial cold yet produced has not exceeded about 220° below zero F.; but it would be rash to assume that it is impossible to produce greater cold. Indeed, it is not improbable that means may yet be found to produce cold sufficient to liquefy either oxygen or hydrogen at ordinary pressures. The question at once arises, what has already been done in the production of extremely low temperatures, and in what direction can such efforts be extended. But practice is dependent upon principles, and for principles we must fall back upon the mechanical theory of heat, thoroughly established and accepted universally by physicists, but as yet imperfectly understood by numbers of chemists educated in the old school of Black and Leslie.

It is useful in the outset to recal certain experiments of Dr. Joule; and, parenthetically, it may be observed that it is to be regretted that no collected edition of the papers of this able man has yet been published. In his paper read to the Royal Society in June, 1844, some very interesting experiments with compressed air are detailed. The paper commences by reference to the well-known fact that compressed air when expanded becomes cooled, that is, when it escapes into the atmosphere; but Dr. Joule asked himself this question—"Would the cooling effect be greater or less were the same air expanded into a vacuum?" He

answered the question by experiment. Two receivers were taken; one was filled with air compressed to about 20 atmospheres; the other receiver was exhausted by an air-pump. The two receivers having been connected by a short pipe in which there was a stopcock, the entire apparatus was immersed in a canful of water. Then by opening the stopcock the compressed air was permitted to flow from the full to the empty receiver.

The advocates of the old theory of heat would have said that since the compressed air in this operation dilated to twice its original volume heat must disappear to meet the increased capacity of air in consequence of its expansion. But the result, indeed, entirely contradicted this. There was no cooling effect whatever measurable by the most delicate thermometer, a result which certainly would have occurred had the compressed air been made to displace the atmosphere. In fact it has since been abundantly proved that the reduction of temperature which a compressed gas undergoes when allowed to expand into the atmosphere is owing to work done in displacing the atmosphere, which offers the well-known resistance of about 15 lbs. per sq. inch to the escape of the gas.

Putting the facts into a brief statement, it may be taken as proved that when a compressed air or gas is allowed to expand the greater the resistance, and consequently the work done in overcoming such resistance, the greater the resultant cooling effect produced. Expand a compressed gas into a vacuum, no cold is produced; expand it in opposing the atmosphere, some cold is produced; expand it against a resistance such as that of a heavily loaded piston so as to contribute to its motion, and the maximum cooling effect is produced. Sir W. Thomson, Rankine, and Clausius have worked out the whole subject mathematically, but it may be convenient to quote Clausius. He takes typical cases to illustrate the difference on the one hand of the expansion of a compressed gas as against the resistance of the atmosphere; on the other hand, as against the resistance of a loaded piston. Assuming an initial pressure of 5 atmospheres in each case, he shows that to bring the temperature of the expanded gas to the initial temperature before expansion, the first case would require 17 units of heat as against 74.9 required in the second case.

Long before the mechanical theory of heat was enunciated Dr. Gorrie utilised for freezing-machines the intense cold produced by expanding a gas behind a working piston. Though his theoretical ideas were somewhat confused, Charles Randolph noticed and commented upon it in the case of the compressed air-engine erected at Govan Colliery in 1849. Air compressed to 20 to 30 lbs. to the square inch was sent down a shaft 176 yards deep, and along a road about 700 yards long, where it was used for working an old steam-engine in the place of steam, the result being that the ports of the engine cylinder were frequently blocked up with ice.

The same phenomena can be observed in any kind of machinery worked with compressed air. In fact I have frequently observed the temperature of air issuing from the cylinder of coal- and rock-cutting machines to register fully many degrees below zero. Indeed, by such a process an unlimited reduction of temperature can be produced, provided the air or gas be subjected to sufficient compression before expansion, and supposing always that care be taken to remove the heat of compression by passing the compressed air through a surface condenser, kept cool with water, or by the actual injection of water at the time of its being compressed. When I first turned attention to the subject I exercised myself considerably in efforts to contrive a method of expanding air or gas so as to cause it to do the maximum of work in the act of expansion. But I speedily became convinced of the unlikelihood of anyone being able to invent a more perfect instrument for converting the expansion of an elastic fluid into work than is afforded by a well constructed steam-engine. Mallard shows us that compressed air made to do work in an engine, at an initial pressure of 10 atmo-

* A paper read to the Chemical Section of the Philosophical Society of Glasgow and to the Institute of Engineers (Scotland). Full drawings of the machinery will be found in the *Transactions* of the latter body, published in Glasgow.

spheres and temperature of 62° , should become reduced to -103° F. on being discharged, putting aside deductions for friction, &c. This illustrates the limitless production of cold by such a process; for let the initial pressure be higher, say 20 or 30 atmospheres, and the cooling effect will correspondingly be produced. Indeed, it seems to me that if Messrs. Pictet and Cailletet had in their experiments made the compressed oxygen or hydrogen do more work in expanding than the mere displacement of the atmosphere, then they would have been liquefied at much lower pressures. M. Pictet employed no less than 9000 lbs. pressure per square inch in his experiments. But it can be shown that a compression of only one-tenth of this, or 900 lbs. per square inch, raises a gas to 1200° F.; and that if gas so compressed be brought down to 60° , and then expanded in the act of doing work, it must necessarily be cooled down to a lower degree than any yet experimentally shown, and which would probably be sufficient to effect the liquefaction of any gas we are acquainted with.

Again, in their experiments there must have been a production of heat militating against the liquefaction from the friction of the gases in escaping into the atmosphere through a minute orifice.

This is the theoretical aspect of the case, but in practice several considerations arise.

It is very difficult constructing the necessary apparatus strong enough without having such a mass of metals as to communicate heat rapidly to its contents; and, further, the friction of the piston rings and the rubbing surfaces of the valves, piston-rods, &c., is difficult to neutralise, lubrication at excessively low temperatures being troublesome.

How far these difficulties may be overcome on original research remains to be proved; but they have been dealt with to some extent in machinery constructed under my direction for technical purposes, and it may be observed that this machinery differs essentially from apparatus previously used, inasmuch as its construction admits of the liquefied gases being produced in a *continuous stream*, whereas in all former experiments with the liquefaction of gases, a small volume has been first confined and liquefied, the experiment then coming to an abrupt end.

In a paper read to the Chemical Society of London (Sept., 1875) I described the effect of pressure and cold upon the gaseous products of the distillation of carbonaceous shales, which remain permanent after all the condensable vapours within are removed in the usual way by cooling to atmospheric temperature. These gases, whilst being permanent at 50° , yield under the influence of a pressure of about 10 atmospheres and a cold of 200° , large quantities of volatile liquid hydrocarbons, consisting chiefly of butylen, amylen, hydride of amyl, and hexylen.

The machine which is the subject of this paper was designed to deal with 300,000 cubic feet per day of this waste gas, by subjecting it to the cold and pressure just mentioned, and has now been working three years, having liquefied, in round numbers, about a quarter of a million gallons of these liquid hydrocarbons. It involves—

- (1.) The pumping of the gas by steam-power into a system of tubes capable of being externally cooled, and from which condensed liquids can be drawn off by ball-cocks.
- (2.) Employing the compressed gas after being deprived of its liquid for working a second engine coupled with and parallel to the first, thus receiving a portion of the force originally employed in the compression.
- (3.) Employing the expanded gas after having had its temperature reduced in the act of doing the work of pumping for supplying the necessary cold for cooling a portion of the condenser pipes to zero.

The machinery is kept in motion by the piston's cylinders, A and A1, the pistons of which are connected with a common crank shaft, both being of 30-inch stroke.

Between these two motive cylinders is a heavy fly-wheel of about 7 tons weight. It became very evident in the early stages of construction that some kind of automatic regulator would be required, so that the speed of the machine should be such as to exercise no undue suction upon the shale retorts, otherwise atmospheric air might have been taken into the machine in admixture with the hydrocarbon gases. This was accomplished by putting a small gasometer governor between the two motive cylinders just described, and adjusted so delicately that a variation of pressure equivalent to 0.25 would, by a system of levers, actuate a throttle-valve and open or close the admission of steam to the cylinder, and thus regulate the speed of the engine. These levers, when once adjusted, cause the engine to run at full speed of 80 revolutions when the pressure in the retorts rose to 0.75 (or $\frac{3}{4}$) inch, and to run at half speed when the pressure sank to 0.25 inch.

We will now trace the progress of the hydrocarbon gases. They are sucked into two pumps worked by the motive cylinders A and A1, the piston rods of which extend to the two pumps, and are therein compressed to about 8 atmospheres absolute, or a little more than 100 lbs. per square inch above the ordinary atmosphere. Under these circumstances some hundreds of degrees of heat are generated which require to be removed: this is accomplished in great part by injecting about 8 cubic inches of water into each pump every stroke of the engine. This water is forced into the gas-pump by small injection-pumps. The compressed gas is now conveyed by the pipes with injected water to the top of a surface condenser, and passes downwards through 200 iron tubes of about 1 inch diameter and 4 feet 6 inches long, these tubes being surrounded with cold water. The mixture of compressed gas, liquefied hydrocarbons, and injected water arrives at length at about 70° F. in reservoir at bottom of the surface condenser. The water and liquefied hydrocarbons are allowed to escape by an automatic regulated cock, and the compressed air is carried to the top of a second surface condenser, precisely similar in construction and with precisely similar outlets at the bottom reservoirs. It is then carried forward to the top of the third surface condenser, where it first passes downwards through a chamber packed with salt, so as to convert any suspended mixture into brine, and afterwards further downwards through 200 more 1-inch iron pipes in the case *not* surrounded with water. The compressed gas-still of about 8 atmospheres pressure is finally used for working the motive cylinder, which does not differ in construction from a steam cylinder, from which it is discharged after doing work considerably below zero Fahr. This constant stream of ejected and cold gas is made to circulate around the condensing pipes of the third surface condenser, and is finally carried away to be burnt as fuel under the boiler, being chiefly a residue of marsh-gas and hydrogen.

The cycle is thus complete, and it only becomes necessary to indicate all the cylinders and note the volumes and temperatures of the incoming water and gas and their final temperatures to have most interesting proof of some of the most important fundamental laws of thermo-dynamics.

The machine was driven by steam of 50 lbs. initial pressure, and when it became desirable to indicate the cylinders the working pressure was kept as nearly as possible 100 lbs. to the square inch, and the piston speed moderate, viz., 144 feet per minute (48 revolutions). Under these circumstances it was found that the—

Motive cylinder A indicated	33 h.p.
" " A1 "	7 "
Total driving power	40 "
The gas-pumps offered a resistance of	32 "
Difference	8 "

The indicator cards show that the volume of compressed gas introduced into cylinder A1 became reduced to atmospheric pressure by the end of the stroke.

It is admitted to the cylinder not at the full pressure of 100 lbs., but by means of a reducing valve at about 60 lbs., being thus cut off at one fourth of the stroke. The recovered power seems comparatively small, but it must be taken into account that there is some loss of gas during its progress through the machine at the points where the liquid hydrocarbons are drawn off (probably 10 per cent); also, that there will be some of the gas liquefied, and that the residue of hydrogen and marsh-gas, being so very mobile, might have a tendency to pass between the piston block rings and the inside face of the cylinders. Taking these facts into consideration, the working of this cylinder has been very remarkable. It is generally coated externally with some inches thick of ice from the freezing mixture deposited upon it from the external atmosphere.

The indicator cards of the gas-pump are very beautiful, and show a very near approach to the true isothermal curve of compression, resulting from the law of Mariotte. No doubt much of the beauty of this card is owing to the extremely small clearance spaces at the end of the stroke of the pump piston, viz., 0.25 of an inch, which again gets filled up with water from the injecting pumps. Such good curves also demonstrate the efficiency of the injection of the cold water by the small force-pumps, and the superfluity of elaborate arrangements, for throwing in pulverised water—insisted upon by Colladen, of Geneva, and other Continental engineers. This machinery has been frequently run at pressures approaching 150 lbs. to the square inch, and piston speed of 240 feet, without ever causing the pump-barrels or valves to reach a temperature of above blood-heat.

The pump-valves are somewhat peculiar in construction, the arrangement being suggested by my friend, Mr. John Thompson, of Messrs. R. Laidlaw and Sons, the builders of the machine. Each pump end has four suction-valves of 1½ ins. diameter, and three delivery-valves of 1¼ ins. diameter. Each of these valves work in a wrought-iron barrel, which contains the spring and the valve-spindle suitably guided. These wrought-iron barrels containing the valves are about 2¼ ins. diameter, and are externally grooved with a thread, so as to enable them to be screwed into the ends of the pump, or with the whole contents removed at pleasure.

The valves have worked very well, but the springs being light require replacing at frequent intervals. Subsequent experience has certainly induced me to prefer valves fewer in number, and of larger diameter so as to admit of stronger springs, but on the whole the arrangement has been very effective.

The piston blocks of the gas-pumps were originally provided with three-eighths square steel rings, but latterly it has been found that the common ¼-inch square cast-iron ring wears best, and keeps the pumps in good working order for fully six months.

The machine has been working continuously from Monday mornings to the following Saturday night as a rule without stopping, and has not been stopped for overhauling or repairs more frequently than the average of steam-engine machinery.

Passing from mechanical details, as an example of results obtained by the machinery the following summary of consecutive observations made over a period of thirty days at the latter end of 1876 may be stated:—

Average temperature of the gas after being expanded and discharged from the motive cylinder A 1	30° below zero F.
Average temperature of the compressed gas on its way to be expanded after leaving the surface condenser S 3	20° above zero F.
Minimum temperature attained by the gas discharged from the motor cylinder H 1	47° below zero F.

Practical considerations as to wear and tear, steam consumption, &c., made it undesirable at that time to increase the working pressure above 135 lbs. to the square inch,

but there is no doubt even lower temperatures would have resulted if the capabilities of the machine had been stretched to their utmost limit.

The hydrocarbon liquids obtained by this process are those produced by pressure alone escaping from the first and second, and those escaping from the third surface condenser at temperatures approaching zero, the result of combined cold and pressure. The former has generally a specific gravity of 0.710 at 60°, the latter of about 0.670 at 60°. The total produce of condensed hydrocarbons has been about 2000 gallons per week, which, when distilled, yield a considerable percentage of extremely volatile liquid hydrocarbons of specific gravities varying from 640° to 0.660. These liquids, judging from the action of bromine, are chiefly amylene, hexylene, and dissolved butylene, with a small admixture of the corresponding hydrides, whereas the extremely light portion of American petroleum are in main composed of the hydrides of the alcohol radicles.

The machinery I have just described was built for Young's Paraffin Light and Mineral Oil Co., for dealing with a portion of the gas produced at their works, probably about one-fifth. There are some who believe that the ends attained by this machine are as well accomplished by processes dependent upon gaseous absorption, the absorbent liquids being oils of heavy specific gravity; but my object in this paper is not to discuss this question, but to call attention to the process as the first application of Faraday's principle of the liquefaction of gases combined with an application of the principles of the mechanical production of cold, first foreshadowed by Count Rumford and Sir Humphry Davy, and developed by Joule, Clausius, Rankine, and Sir W. Thomson.

IMPROVED METHOD FOR MAKING PLATINUM-ALLOY ASSAYS.

By NELSON W. PERRY, E.M.

ABOUT a year ago I had occasion to make some assays of platinum alloys. The first method employed consisted of cupellation with addition of Ag and Pb, the loss being base metal. The silver was obtained by precipitating from solution in HNO₃ with HCl. This necessitated bringing the precipitate on a filter, washing thoroughly (giving a large filtrate for evaporation), drying, weighing, and calculating the Ag from weight of AgCl obtained. I will not detail this method further, but mention this last step to show one of the tedious steps in the operation. Doubtless many of your readers are familiar with this assay. The results by this method—strange to say, an established one—I found exceedingly unsatisfactory and inaccurate; one of the causes of its inaccuracy being that the percentage of platinum is obtained by difference, namely, all the other constituents being determined, their combined percentage is subtracted from 100. The error is therefore cumulative for platinum. The assay, or rather combined assay and analysis, as it is, required almost as much time as a good quantitative analysis, and yet could lay no claim to equal accuracy.

I therefore set myself to work to devise, if possible, some better method, the results of which I set before you.

In platinum alloys, or native platinum, the metals to be determined are base metal, Ag, Au, Pt, and iridosmine.

Base metal is removed from the others by cupellation. Silver is soluble in H₂SO₄, while the others are untouched.

Platinum when alloyed with twelve times its weight of silver, is soluble in HNO₃.

Gold is soluble in aqua regia, and iridosmine untouched by acids.

Making use of the above properties, I was enabled to effect a separation of the metals both accurately and rapidly, no filtering being required, and all the washing being done by decantation.

Assay of platinum alloys containing base metal, silver, platinum, gold, and iridosmine.

Charge Pt 200 Mg, pure Ag 150 Mg, or sufficient to produce perfect cupellation (exact weight.)

Wrap charge in sheet lead and cupel. Weigh button. Loss = base metal.

Flatten button, anneal, roll out thin, anneal again, and make into cornet as in gold bullion assay. Introduce cornet into parting flask and part with concentrated H_2SO_4 . Wash, anneal, and weigh. Loss from previous weight = Ag in original alloy + Ag added for cupellation.

Alloy cornet with at least twelve times the amount of Ag that there is Pt present, and, as before, form cornet and part first with HNO_3 sp. gr. 1.16, and then HNO_3 sp. gr. 1.26. Wash thoroughly, anneal in annealing cup, and weigh. Loss = Pt.

Treat residue with *aqua regia*, obtain Au by loss—the residue is iridosmine.

Time to complete assay in duplicate, 2 hours 45 minutes.

The quality of Ag added should at least be sufficient, so that after the addition the Ag in the alloy will be to the Au as 3 : 1.

As platinum and iridosmine add greatly to the infusibility of the compound, silver in sufficient quantity must be added to prevent "freezing" and give a perfect cupellation. Any large excess over these requirements is to be avoided—first, because the residue, after parting, will in that case be non-adherent and in a more or less fine state of subdivision, which may occasion loss in washing by decantation; second, the larger the button cupelled the more difficult it is to obtain a good cupellation, and the greater the loss of Ag during the process. It may, for this reason, sometimes be necessary to use only 100 Mg of the alloy for assay instead of 200 Mg, as above.

The cupellation should take place at a moderate temperature, until near the "blick," when the assay should be thrust back into the hottest part of the furnace to prevent "freezing." The button must remain in the muffle until all the Pb is gone.

In parting with H_2SO_4 , boil for several minutes. In other respects, this operation is identical with the gold-bullion assay. Any large excess of Ag over twelve times the amount of Pt in alloy is to be avoided, as it causes the residue, after parting, to be too fine and float, thereby occasioning loss in washing. Insufficient Ag is even worse, as the Pt will then be only incompletely dissolved.

By the above method I was successful in obtaining very closely duplicating results, was enabled to do away with all precipitation and filtering, no apparatus being required, except that necessary for the gold and silver bullion assay, and, what is only second in importance to accuracy, attained results with the least expenditure of time. Mr. William Strieby, A.M., E.M., has assisted me materially in perfecting this assay, and employed it in his own work with equally satisfactory results.

I present the above to your readers, hoping it may prove of service to some engaged in similar work.—*Engineering and Mining Journal*.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, February 20, 1879.

Dr. J. H. GLADSTONE, F.R.S., President, in the Chair.

AFTER the confirmation of minutes, &c., the following certificates were read for the first time:—W. North, F. Podmore, W. Y. Gent, W. Radford.

The list of Officers for the ensuing year as proposed by the Council was then read —

President—Warren de la Rue, F.R.S.

Vice-Presidents—The list remains unaltered with the insertion of J. H. Gladstone and omission of Warren de la Rue.

Treasurer—W. J. Russell.

Secretaries—W. H. Perkin, H. E. Armstrong.

Foreign Secretary—Hugo Müller.

Other Members of the Council—M. Carteghe, A. H. Church, W. N. Hartley, C. W. Heaton, E. Riley, W. Chandler Roberts, T. E. Thorpe, W. Thorp, J. L. W. Thudichum, W. A. Tilden, R. V. Tuson, R. Warrington.

The following gentlemen were elected auditors for the year:—R. J. Friswell, J. Spiller, and J. M. Thomson.

The PRESIDENT then called on Dr. OTTO N. WITT to read a paper "On Colouring Matters Derived from Diazo-compounds." During the last three or four years aniline colours have come into such general use as to, in many instances, replace the old costly artificial and natural dyes. Although there has been known for some time a great variety of magenta, violet, and blue aniline dyes, green and yellow dyes were almost unknown until quite recently. Three years ago the author, in a paper read before the Society, indicated a theory as to the relation between the chemical constitution and the colouring power of aromatic substances. In this paper he pointed out that there existed a series of compounds, the colour of which was in close relation to their chemical constitution. The first link of this series is azo-benzene, $C_6H_5-N=N-C_6H_5$. This substance is of a deep yellow colour, but being devoid of salt forming groups is not applicable for dyeing. By introducing amido- or oxy-groups, compounds can be formed having strong affinities for textile fibres, which increase with the addition of each salt-forming group into the molecule. At the time this paper was read amido-azo-benzene and tri-amido-azo-benzene were the only dyes of this class known; the first was of a beautiful bright yellow colour, but fugitive; the second was fast, but dull. Since that time the intermediate di-amido-azo-benzene has been formed and named chrysoidin, which combines the beauty of the monamido- and the fastness of the tri-amido-bodies. The preparation of chrysoidin is then given. Griess has shown that not only can amido-azo-compounds be obtained, but that the oxy-derivatives of azo-benzene and other similar bodies can be prepared by the action of diazo-compounds on the corresponding phenols; these oxy-derivatives are also dyes; so that these di-azo-compounds have opened out an almost inexhaustible mine of new and beautiful dyes. Since Hofmann's paper on chrysoidin investigations on this class of compounds have been set on foot in the laboratories of almost all aniline colour manufacturers. The patents taken out for their production are constantly increasing in number. In consequence of his connection with Messrs. Williams, Thomas, and Dower the author has not been able to carry out his original intention of describing fully the preparation and properties of all his new azo-colours. The prototype of these compounds is chrysoidin. The author has already given a description of the properties of this substance in *Journ. Chem. Soc.*, ii., 457, 1877. He has since prepared in a pure state the following analogous substances closely resembling the typical product:—Ortho-tolyl-phenylen-chrysoidin, para-tolyl-phenylen-chrysoidin, phenyl-toluylen-chrysoidin, ortho-toluylen-chrysoidin, para-tolyl-toluylen-chrysoidin, phenyl-toluylen-chrysoidin sulphonic acid, α -naphthyl-phenylen-chrysoidin sulphonic acid, α -naphthyl-toluylen-chrysoidin sulphonic acid. Each basic colour has an acid counterpart similar in shade and constitution but containing hydroxyl- in place of amido-groups. The acid counterpart of chrysoidin has been prepared by Baeyer and Fägar, and studied by Typke. This compound is a beautiful but unstable dye. The author therefore introduced a sulpho-group into its molecule by treating resorcin with para-di-azo-benzene sulphonic acid. The substance has since been described by Griess. Its constitution that meta-di-azo-oxy-benzene sulphonic

acid. Its acid sodium salt is sold under the name of tropæolin O. By a similar reaction monoxo-azo-benzene sulphonic acid has been obtained from phenol. Its sodium salt is known as tropæolin Y. Tropæolin OOO No. 1 is oxy- α -naphthyl, and tropæolin OOO No. 2 the sodium salt of oxy- β -naphthyl-azo-phenyl sulphonic acid. Tropæolin OOOO is isomeric with tropæolin OOO Nos. 1 and 2. Another class of azo-colours can be obtained if the salt-forming properties of amido-azo-benzene and analogous compounds be intensified by introducing one or more sulpho-groups. Thus, by acting with para-di-azo-benzene sulphonic acid on dimethyl-anilin, a dye is obtained, in which, however, the basic properties are too prominent, and to obtain a proper equilibrium one phenyl instead of two methyl groups must be introduced into the molecule of amido-azo-benzene sulphonic acid. Thus is produced one of the most beautiful of the azo-colouring matters. It is known as tropæolin OO. The author gives a detailed description of the preparation and purification of phenyl-amido-azo-benzene, which, when pure, forms leaflets or needles of a fine golden yellow colour; M.P. 82° ; soluble in benzolin, alcohol, ether, and benzene. The author reserves for a future communication a description of amido-diphenyl-amin. Phenyl-amido-azo-benzene when treated with amyllic nitrite and acetic acid yields a nitrosamin crystallising in orange needles, melting at 119.5° ; its formula is $C_{18}H_{14}N_4O$. By the action of di-azo-benzene-sulphonic acid on diphenyl amin, tropæolin OO is obtained. It is a powerful acid and forms well-defined salts. The author gives a description of the potassium, sodium, ammonium, trimethylamin, barium, calcium, and aniline salts. In conclusion the author trusts that he has succeeded in giving a sketch of what may be called the genuine azo-colours, the true oxy- and amido-derivatives of azo-benzene, and analogous compounds. Compounds derived from amido-azo-bodies by the action of amines as well as coloured substances containing the azo-group, $-N=N-$, may also be termed azo-colours. The author hopes in a future paper to lay before the Society his researches on these more intricate compounds.

The PRESIDENT said that all must have appreciated the very lucid and brief manner in which Dr. Witt had described this beautiful series of compounds, interesting both from a scientific point of view and from their application as dyes.

Mr. PERKIN said that the paper was of peculiar interest to him, as showing the enormous strides which aniline dyestuffs had taken during the last few years. He would like to ask whether these dyes were suitable for silk, wool, and cotton?

Dr. ARMSTRONG remarked that the brief description given by Dr. Witt conveyed but a slight notion of the enormous amount of work concentrated in the results. He would ask Dr. Witt if he could point out the influence of the different groups in modifying colour? and also if he had any experience of tropæolin OO as an indicator in alkalimetry?

Dr. WITT said that most of the dyes, especially the sulpho-compounds, were only suitable for silk and wool, the amine compounds dyed cotton. He had already pointed out two groups concerned in the tinctorial power of a substance, which he had called chromogens and chromophors (*Chem. Journ. Soc.*, ii., 403, 1876). Subsequent results had confirmed these conclusions. Oxy- and amido-groups principally had to do with the colour of the dyestuff. The sulpho-group weakened the colour but conferred stability on it. Also, *ceteris paribus*, the more oxy- or amido-groups contained the more powerful is the dye. It does not follow, however, that the beauty of the colour increases with its intensity. The heavier the molecule the more the colour tends to the violet end of the spectrum. Some importance must be attached to the localisation of the salt-forming groups, i.e., the nearer they are together in the molecule the more powerful is the colour. Tropæolin OO has been recommended as an indicator by Von Mütler. It is not affected by CO_2 , and only after

some time by SH_2 , so that it can be used to titrate crude soda. A still better indicator is the dimethyl-diazo-benzene sulphionate of ammonium.

Dr. RUSSELL then took the chair during the reading of a paper by Dr. GLADSTONE entitled, "*Investigations into the Action of Substances in the Nascent and Occluded Conditions (Hydrogen, continued)*," by J. H. GLADSTONE and A. TRIBE. From a recent study of the behaviour of nascent and occluded hydrogen (*Chem. Soc. Journ.*, 305, 1878) the authors concluded that these conditions of the element are not, as hitherto supposed, different, but are closely related if not identical, and that the activity of the so-called nascent hydrogen is the consequence of its intimate association with the metals employed to bring about the liberation of the element. In the present paper the authors have examined the action of nascent and occluded hydrogen on nitric and sulphuric acids. The nascent hydrogen was obtained by electrolytic decomposition of the acids in Hofmann's arrangement for illustrating the composition of water electrolytically. Nascent Hydrogen and Nitric Acid:—Faraday and Bourgoine have shown that electrolytic hydrogen reduces strong nitric acid, but only imperfectly or not at all when diluted with an equal bulk or more of water. If the oxidation of the freed hydrogen in this action results from its being in the occluded condition, the reduction of the acid would depend on its strength only in so far as this facilitated the de-occlusion of the hydrogenised electrode, and the stronger acid might be expected to do this the more readily. And it follows that with a given strength of acid the amount of free gaseous hydrogen should bear some relation to the rate at which the electrolysis takes place, for were the gas freed from its nitric radical, not faster than it could be occluded, none should pass through the liquid; but if the evolution of the gas were faster than the occlusion, free hydrogen should escape. The authors have proved that this is the case. Their results are given in the annexed table—

Grove's Cells Used.	Proportion of Acid to Water.	Time of Experiment.	Reduction in c.c. of Oxygen.	Free H.
1	1 acid	3 hours	31.5	0
2	68.2	55 mins.	28.4	0
4	per cent	22 "	28.8	0
8	0 water	10 "	29.4	0
1	1 acid	215 "	30.6	0
2	to	42 "	29.0	2 c.c.
4	1 water	19 "	24.2	9 "
8		8 "	16.8	22 "
2	1 acid	61 "	17.0	30 "
4	to	18 "	0.5	65.7
8	2 water	8 "	0.3	68.7

Each experiment went on till 35 cc. of oxygen collected at the anode. In the 2nd, 3rd, and 4th experiments of the series 1 to 1 the evolution of hydrogen at the cathode ceased quite suddenly at the end of three minutes. This is due to the presence of nitrous acid, which prevents the escape of hydrogen with eight cells when present in the proportion of 0.059 grm. to 100 cc. of 1 to 1 acid. Occluded Hydrogen and Nitric Acid:—Nothing is known as to the action of occluded hydrogen on nitric acid. Dr. Armstrong infers (*Chem. Soc. Journ.*, 1877, 82) that it has no action. The authors charged some finely-divided platinum with hydrogen. On pouring on it some pure nitric acid the metal became red-hot, the liquid became yellow, and nitrous fumes escaped; so that occluded as well as nascent hydrogen acts on nitric acid. Palladium charged with hydrogen dissolves in nitric acid 1 to 1 without setting free any gas, so that the hydrogen in this case must be oxidised. Nascent Hydrogen and Sulphuric Acid:—The acid was decomposed electrolytically. The authors conclude that hydrogen associated with platinum reduces oil of vitriol very readily, sulphurous acid being sometimes formed; a film of sulphur also appears on the negative electrode. Occluded Hydrogen and Sulphuric

Acid:—Finely divided platinum did not reduce the acid, but the metal charged with hydrogen immediately gave the odour of sulphurous acid when the sulphuric acid was poured on it. Platinum charged with hydrogen also reduces sulphuric acid. In conclusion, the authors give the results of some experiments made by plunging a small piece of sheet magnesium into an excess of strong nitric acid diluted with an equal bulk of water. The metal dissolved in one or two seconds, gas being given off which was combustible and explosive and contained hydrogen. The authors claim to have demonstrated the possibility of the replacement of hydrogen in nitric acid by a metal, and to have established the close likeness of character, and therefore of condition, between the so-called nascent hydrogen and the hydrogen occluded by metals.

Dr. ARMSTRONG said that he had not stated in the paper referred to by Dr. Gladstone that occluded hydrogen had no action on nitric acid, but had asked the question has occluded hydrogen, &c.? He had dissolved some pure nickel, given to him by Dr. Russell, in nitric acid and obtained hydrogen, whereas crude nickel gave none. He therefore concluded that the pure nickel might contain some occluded hydrogen. Though admitting the extreme value of the authors experiments, he did not quite agree with all their conclusions.

Dr. RUSSELL said that he had heated and pumped the nickel referred to repeatedly, but had never succeeded in obtaining any hydrogen.

Dr. WRIGHT suggested that the metal might contain a trace of oxide, and when heated, the hydrogen would be converted into steam.

Dr. GLADSTONE briefly replied and then took the chair.

Mr. J. T. BROWN then read a short paper "*On some Methods of Vapour-Density Determinations.*" The author criticises the methods and especially the formulæ of Hofmann, Wertheim, W. M. Watts, Goldschmidt, Freichs, and Meyer. As the actual tensions of the vapour of mercury under various conditions are not known the author suggests that they might be determined by estimating the vapour-tension of a substance over Wood's metal and mercury at different temperatures. He offers the suggestion in the hope that some one will take up the subject. Wood's metal is a fusible alloy, composed of Bi₁₅, Pb₈ Sn₄ Cd₃.

Drs. WITT and ARMSTRONG pointed out that the new method suggested by Meyer was almost perfect, as exact results could be obtained by it with facility.

The SECRETARY then read a paper "*On the Decomposition-Products of Quinine and the Allied Alkaloids.*" by J. J. DOBBIE and W. RAMSAY. In a previous paper the authors gave the results of their experiments on the oxidation of quinine by permanganate. In the present paper the authors have extended their investigation to the oxidation-products of quinidin, cinchonin, and cinchonidin. All these bodies yield, by oxidation with permanganate, acids which are physically and chemically identical. This acid, the authors prove by analysis, &c., to be tricarbo-pyridenic acid. They give in detail the method employed. More than 10 per cent of the acid was obtained from each base. The paper contains an account of the properties and form of crystallisation of the acid, with many analyses. The acid is tribasic. Potash, soda, ammonia, silver, calcium, barium, strontium, zinc, and copper salts were prepared, and are described. The formula deduced from the analysis of the acid and its salt is $C_8H_5NO_6 + 1\frac{1}{2}H_2O$. The result of the present investigations confirms the conclusions previously arrived at by the authors, viz., that there is a close relation between the cinchona bark alkaloids and the bases of the pyridin series, and proves that the four principal alkaloids derived from the cinchona bark all yield on oxidation the same acid. In conclusion the authors draw attention to the fact that their first paper was published March, 1878, and that in the *Ber. der Deut. Chem. Gesell.*, Feb. 11, 1879, is a paper by Hoogeweff and Von Dorp, which confirms the authors researches as regards quinine.

The Society then adjourned to March 6, when the following papers will be read:—"The Quantitative Blowpipe Assay of Mercury," by G. Attwood; "On Gas Analysis and Gas Apparatus," by J. W. Thomas; "The Isomeric Dinaphthyls," by Watson Smith; "On the Action of Isomorphous Salts in Exciting the Crystallisation of Super-saturated Solutions of each other," by J. M. Thomson.

PHYSICAL SOCIETY.

Ordinary Meeting, February 22, 1879.

Prof. W. G. ADAMS, President, in the Chair.

New Members:—Rev. Coutts Trotter, Prof. G. D. Liveing, J. C. Adams, F. W. Paterson.

Dr. C. SIEMENS described his New Electric Current Regulator. A necessary condition of the transmission of power to a distance by electricity along a single conductor and re-distributing it by means of branch circuits to separate electric lamps or motors, is that the current strength in each lamp shall be practically uniform; otherwise the current flowing in the whole branch. Hence the necessity of a regulator to regulate the flow of current so as to keep it uniform, however the resistance of the circuit, or the electromotive force of the source, may vary. The author believes that by properly arranging a number of dynamo-electric machines, either in series or parallel (for intensity or quantity), at each end of the wire, a vast amount of power may be sent along a small copper conductor successfully, provided the distribution is properly regulated. He has designed a regulator based on the heating of a wire by the passage of a current through it. A fine strip of mild steel $\frac{1}{16}$ m.m. thick is stretched horizontally between two terminals. An upright spindle is supported by means of an insulating foot upon the middle of this strip in such a manner that as the strip bends or sags by its expansion, the spindle sinks with it. Now this spindle carries at its top a table or plate of metal (or, as the case may be, a set of radial springs), and as the spindle rises or sinks to different heights this plate or these springs make contact with other springs set radially round; and these contacts take out from or throw in resistance coils into the circuit of the current. The sensitive strip is so thin that it may be regarded as a radiating surface merely, and it may be assumed that its temperature, due to heating by the current, balances itself with the radiation instantaneously. After passing through the steel strip the current flows through the coils thrown into circuit, and, by the arrangement we have described, if the current increase so as to overheat the strip, the latter sags a little more, the spindle sinks, and the consequence is that one or more of the spring contacts is broken, and one or more coils inserted in circuit. A rise of 1° F. in the temperature of the strip is sufficient to liberate two or three of these coils. The fact that the temperature of the strip varies as the square of the current favours the sensibility of the apparatus. An older form of this apparatus having pendulous contacts was also shown; also a regulator in which the expansion by heating of a sensitive wire caused the resistance of several carbon buttons in contact to vary through the pressure exerted on them by means of a bell cranks lever. Dr. Siemens had not been able to prepare carbons which gave the wide variations of resistance obtained by Mr. Edison. Siemens's regulator can also be used as a current meter by causing the sensitive strip to actuate a lever, carrying at its end a pencil writing on a moving paper.

Dr. COFFIN said that he had thought of a regulator in which the heating of a wire spiral in a gaseous chamber would cause the gas to expand and drive up a mercury column past a series of contacts, which would throw resistances in circuit.

Dr. GUTHRIE suspected from some experiments of his

that the conductivity of metals was not strictly proportional to their sectional area.

Dr. SCHUSTER then gave the results of some observations of his on the Spectrum of Lightning. These were made by a spectroscope with two prisms, one for the red and the other for the blue end of the spectrum, which were shifted into the line of sight by a chamber arrangement. Three observations were made, one at Las Ammas, one at Maniton, and one at Salt Lake City, last year. These showed the three nitrogen lines, with three well-defined bands, and one doubtful band. The nitrogen lines correspond to the spectrum of air, and the bands appear to Dr. Schuster to agree with the spectrum of the light round the negative pole of the spark in a tube containing oxygen with adulteration of carbonic oxide.

Prof. AYRTON then exhibited an Exisothermal Model of a Cooling Globe. If we imagine a globe initially heated throughout to a uniform temperature (as was probably the earth), and then kept in a space having a constant temperature but much lower than that to which the globe was heated, then the temperature at every point of the ball will fall, but at very different rates, the parts for example near the surface cooling comparatively rapidly, while those near the centre will cool very slowly. A surface could, therefore, be constructed, such that the x of any point on this surface represented its distance from the centre of the globe, the y the time t from the commencement of the cooling, and the z the temperature of that point at that moment. The nature of this surface would depend on the size of the globe, on the specific heat, conductivity, and surface emissivity of the material. Now the "Experiments on the Heat Conductivity in Stone" of Profs. Ayrton and Perry, described in the *Philosophical Magazine* for April, 1878, enables them to determine these constants accurately for a trachyte sphere, and by using these data they have been enabled subsequently to construct such a surface called by them an "exisothermal" one for a trachyte globe of 8000 miles in diameter, and which gives graphically the temperature of every single point of the earth from the moment when it was at the temperature of molten trachyte down to eight hundred thousand million years afterwards, that is, until long after the present era.

OBITUARY.

PETER LE NEVE FOSTER, M.A.

THE Society of Arts has just sustained a severe and sudden loss by the death of its well-known Secretary, Mr. Peter Le Neve Foster. On Thursday, February 20, immediately on his return to his own house at Wandsworth, Mr. Foster was seized with a sudden attack of heart disease, and some of his family coming into the room, where he had been sitting by himself for a few minutes reading the newspaper, found that he had fallen back from his chair, dead. So little expected was the illness that he had finished his ordinary day's work at his office, and had even walked up from the railway station to his own house.

Mr. Foster was born August 17, 1809, and was the son of Mr. Peter Le Neve Foster, of Lenwade, Norfolk. He was educated under Dr. Valpy, at the Norwich Grammar School, from whence he went up to Trinity Hall, Cambridge. After having taken his degree as Thirty-eighth Wrangler in the Mathematical Tripos of 1830 he was elected Fellow of his College. He was called to the Bar at the Middle Temple in 1836, and practised as a Conveyancer till he became Secretary to the Society of Arts in 1853.

Mr. Foster was intimately associated with all the earlier great exhibitions. He was appointed to carry into effect the provisions of the Act for the Protection of

Inventions in the Exhibition of 1851, and was also named Treasurer for payment of all executive expenses in the original Commission. During his term of office the Society of Arts has flourished as it never previously did, and owing in no small degree to his exertions it has quadrupled its number of members and increased its resources in a still greater proportion. Much of its work was originated by the late Secretary, and all of it was carried out by him.

From his boyhood upwards Mr. Foster took a keen and enlightened interest in many branches of science. He was one of the first to take up and practice, as a scientific amateur, the art of photography, and on this subject he has written a good deal in the pages of the *Photographic* and other periodicals. He was one of the Founders of the Photographic Society, and was on its Council for many years. He was President of the Quekett Microscopical Club for a year, and also served for some time on the Council of the British Association, the meetings of which he has attended regularly for the past twenty years. For many years he acted as Secretary of the Mechanical Section of the Association. He read several papers before the Society of Arts, and was, of course, a constant contributor to its *Journal*, the whole series of which, from the middle of the first volume, was published under his direction.

Mr. Foster leaves behind him a very numerous body of friends, to all of whom his genial and kindly character had endeared him. On the occasion of his completion of twenty-five years' service as Secretary, a strong committee was formed to present Mr. Foster with a testimonial. The list for this was just about to be closed, the amount subscribed being over £1200. Under present circumstances it is probable that a fresh effort will be made to increase this amount, so that a fitting memorial may be presented to Mrs. Foster.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 60, December, 1878.

Report Presented by M. Personne on the Process of MM. Lecourt and Guillemare for Using Chlorophyll in Place of the Copper Salts Employed in the Preparation of Preserved Fruits, Green Vegetables &c.—The patentees find that if the vegetable fibre is placed in contact with soluble chlorophyll it becomes saturated with the colour at about 100°.

Crystallisation of Sugar and the Manufacture of Sugar-candy.—G. Flourens.—A lengthy paper not adapted for useful abstraction.

Biedermann's Central-blatt.

Heft 1, January, 1879.

Comparative Observations of Rain-fall According to Fautrat's Method.—A. Johnen.—The author found the quantity of rain in a beech-wood 13 per cent greater than in the open country. In a pine-wood the excess was only 2 per cent.

Cohesive Power of Various Kinds of Soil.—Prof. F. Haberlandt.—The author moistened different kinds of earth with water, moulded them into cylinders of equal size, and ascertained their "breaking weight" in the usual manner. The cohesive power is affected chiefly by its mechanical structure, not by its degree of fineness.

Every soil has at some degree of moisture a minimum cohesive power.

Heat-capacity of the Constituents of Soils.—Dr. C. Lang.—The specific heat of the several constituents of soils differs much less if referred to volume than to weight.

Manure Experiments on the Agricultural Trial-field of the University of Giessen.—Prof. A. Thaer.—The quantity of nitrogen given to a selected "morgen" of land in the form of manures during seven years amounted to 89.5 lbs. The crops reaped during the same years contained together 198.69 lbs. of nitrogen. Consequently the soil had given 109.19 lbs. nitrogen more than it had received. Hence, whilst the manures furnished 45 per cent of the nitrogen in the crops, the atmosphere had yielded 55 per cent.

Trade Cattle Foods.—Dr. E. Wein, Prof. J. Nessler, Prof. A. Mayer, and Prof. R. Heinrichs.—According to the above chemists the composition of an English cattle food is—

	Wein.	Nessler.	Mayer.
Moisture	13.72	—	12.9
Albuminoids	11.75	13.21	9.4
Oily matter	4.28	4.24	3.6
Non-arotised extractive ..	67.24	—	66.9
Vegetable fibre			3.6
Ash	3.01	—	3.6

Wein and Mayer, from chemical and microscopical evidence, conclude that the food is composed of equal parts of bruised maize and locust-beans, and that the essential oil of a plant of the umbelliferous order has been added. Another English cattle food in the form of cake contained also a large proportion of locust-beans.

Fustus Liebig's Annalen der Chemie,
Band 194, Heft 2 and 3.

Synthesis of Tin-phenyl-compounds.—Dr. B. Aronheim.—This paper contains an account of tin-phenyl-chloride and of its derivatives tin-diphenyl-hydroxyl-chloride, tin-phenyl-oxide, tin-phenyl-dichloride, tin-phenyl-chloro-bromide, tin-phenyl-chloro-iodide, tin-phenyl-dibromide, and of the action of sodium ethylate upon tin-diphenyl-dichloride and the formation of tin-triphenyl-chloride by the action of sodium amalgam or of ammoniacal gas upon tin-phenyl-chloride.

Studies on Phosphates.—E. Erlenmeyer.—A very detailed account of mono-ferro-phosphate, including its behaviour with water and alcohol; of mono-ferri-phosphate, of mono-diferri-phosphate, di-ferri-phosphate, tri-ferri-phosphate, mono-aluminium-phosphate, and tri-aluminium-phosphate.

Contributions to the History of the Naphthalin Series.—J. Stenhouse and C. E. Groves.—The authors describe β -naphtho-quinon, nitro- β -naphtho-quinon, dinaphthyl-diquinhydrin, dinaphthyl-diquinon, dinaphthyl-diquinol, and dinaphthyl-diquinhydrin. The authors have observed that benzo-quinon and α -naphtho-quinon yield diquinons and diquinols on condensation.

Action of Potassic Bibromate with Sulphuric Acid upon Cholic Acid.—Dr. H. Tappeiner.—The products obtained are cholesteric acid, stearic acid, lauric acid, and cholic acid, which are here successively described together with certain of their derivatives.

On Triphenyl-methan and Rosanilin.—E. and O. Fischer.—This valuable treatise is unfortunately not susceptible of useful abstraction.

MISCELLANEOUS.

Gossip in the Provinces.—(From a Correspondent).—The subject of the greatest interest for chemists just now is the new phase of the water-analysis question. Dr. Tidy's paper, read before the Chemical Society, is generally

regarded as a very valuable contribution, containing as it does the results of years of work, and affording chemists another factor of value for judging of the character of drinking-water. There seems indeed too much tendency on the part of the adherents of one or the other system of water-analysis to swear by that method which they are accustomed to use and to pooh-pooh all others as unreliable. Thus Dr. Frankland appears to suppose that chemists who use any other method than his own do so to avoid the trouble of determining the organic carbon and nitrogen. As pointed out by Dr. Dupré, this assumption is unwarrantable, though certainly the facility with which a process may be carried out is an important consideration when there is a choice of alternative methods. There is, however, a very widely spread feeling among chemists that Dr. Frankland has never sufficiently met Mr. Wanklyn's objection respecting the action of nitrates on readily decomposable organic matter. It is clearly not an answer to this objection to quote the figures obtained by the analysis of sulphate of quinine, either in presence or absence of nitrates, though we are unable to recall any experiments to ascertain the effect of the presence of a large excess of nitrates on the determination of the nitrogen of carbon of even such a stable body as the sulphate of quinine.

Dr. Tidy did good work by protesting against the attempt to reduce water-analysis to the empirical application of a few simple tests. On the contrary, the more data the chemist has on which to base his opinion the more correct that opinion is likely to be.

We fear that Mr. Wanklyn has done extensive harm by not sufficiently insisting on the consideration of the supplementary data, and by teaching chemists of limited experience and knowledge of the science to rely too implicitly on the indications given by the ammonia process. We know of chemists who restrict their examination of drinking-waters solely to the ammonia process, and do not even supplement it by a determination of the chlorine and total solids as recommended by Mr. Wanklyn. There is also a widespread habit of ignoring the nitrates and nitrites on the ground that when the oxidation has reached that stage no objectionable impurity can remain. Those who argue thus must be ignorant of the extreme readiness with which ammonia undergoes oxidation in a porous soil, whilst a change of weather, by choking the pores of the soil, will cause a water which previously contained nitrates and no ammonia to present the strongest evidences of the presence of unoxidised sewage.

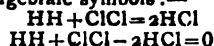
Respecting the determination of the organic nitrogen of carbon of drinking-water, it is interesting to note that distinct advance has been made towards ready means of obtaining these data. Thus Dittmar has shown that accurate determinations of the nitrogen could be obtained by igniting the residue with soda and baryta—not with soda-lime, as stated by Mr. Hartley—and Dupré has quite recently described an easy method of estimating minute quantities of carbon. If these methods bear rigid examination there is a future for the disciples of Dr. Frankland that is very hopeful.

What have chemists been about all these years that they leave it for a metal-broker to utilise the Bessemer process for treating sulphides? Mr. Hollway's paper at the Society of Arts fairly takes our breath away. Fancy blowing a stream of air through molten sulphide of iron for ten consecutive hours, and keeping up the temperature merely by the addition of lumps of raw pyrites. Mr. Hollway appears to have no difficulty in keeping up the temperature, but whether he will succeed in obtaining a fairly rich regulus without material loss of copper in the slag and by volatilisation is another question. One of the most curious points about the process is that more than half the sulphur appears to be liberated in a free state. As pyrites has always hitherto been treated in close vessels at comparatively low temperatures it may very possibly be that a different reaction occurs at a white heat. At any rate the analyses made seem to show that in the regulus

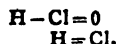
obtained there is often much less sulphur than corresponds to the formula FeS —a fact that points to the existence of a lower sulphide of iron than any hitherto recognised.

The arguments which Mr. Lockyer has laid before chemists in support of his ideas respecting the dissociation of the elements, though ingenious and dealing with highly interesting and suggestive facts, have produced some feeling of disappointment. The definite and somewhat sensational reports which appeared in some of the daily papers had prepared chemists for facts more tangible than those ultimately adduced in support of the theory. Still Mr. Lockyer has got hold of a good idea, and one which will be certain to repay further investigation. Chemists will look forward to future accounts of his progress.

Does Mr. Lockyer know that the identity of many of the chemical elements can be demonstrated mathematically? The following instance, proving the identity of chlorine and hydrogen is, we believe, due to Prof. Clifford, and affords a good illustration of the way in which chemists misapply algebraic symbols:—



Then, extracting the square root from each side of the equation—



and

MEETINGS FOR THE WEEK.

- MONDAY, 3rd.**—Medical, 8.
— Society of Arts, 8. "Dwelling Houses: Their Sanitary Construction and Arrangements," by Dr. W. H. Corfield, M.A. (Cantor Lectures.)
— London Institution, 7.
— Royal Institution, 3. General Monthly Meeting.
TUESDAY, 4th.—Civil Engineers, 8.
— Royal Institution, 3. "Animal Development," Prof. Schäfer.
— Zoological, 8.30.
WEDNESDAY, 5th.—Society of Arts, 8. "The Social Necessity for Popular and Practical Teaching of Sanitary Science," by Joseph J. Pope, M.R.C.S., L.S.A. Pharmaceutical, 8.
THURSDAY, 6th.—Royal, 8.30.
— Royal Institution, 3. "Sound," Prof. Tyndall.
— Royal Society Club, 6.30.
— Chemical, 8. "The Quantitative Blowpipe Assay of Mercury," by G. Attwood. "On Gas Analysis and Gas Apparatus," by J. W. Thomas. "The Isomeric Dinaphthyls," by Watson Smith. "On the Action of Isomorphous Salts in Exciting the Crystallisation of Supersaturated Solutions of Each Other," by J. M. Thomson.
FRIDAY, 7th.—Royal Institution, 9. "Sensation," by Prof. Huxley.
— Geologists' Association, 8.
— Society of Arts, 8. "The Plants of India Adapted for Commercial Purposes," by John R. Jackson.
SATURDAY, 8th.—Royal Institution, 3. "Colbert and Richelieu," by Mr. Walter H. Pollock.
— Physical, 3. "On a New Theory of Terrestrial Magnetism," by Prof. Ayrton and Perry. "On some Experiments with the Quadrant Electrometer," by Dr. J. Hopkinson. "On the Maintenance of Constant Temperatures and Pressures," by F. D. Brown.

TO CORRESPONDENTS.

ERRATUM.—P. 67, col. 2, line 12 from top. The words "I am told the substance was not quite pure" applies only to the chrysene. The dinaphthyl was beautifully white and absolutely pure.

W. E. Robinson and Co.—You had better advertise for a market in our columns. There are plenty of uses for the salt.

J. G. H.—We regret we cannot give the desired information. Our correspondent had better advertise in the *CHEMICAL NEWS*.

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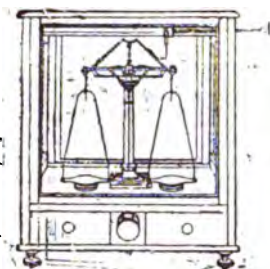
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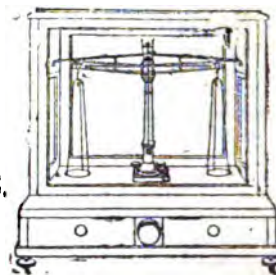


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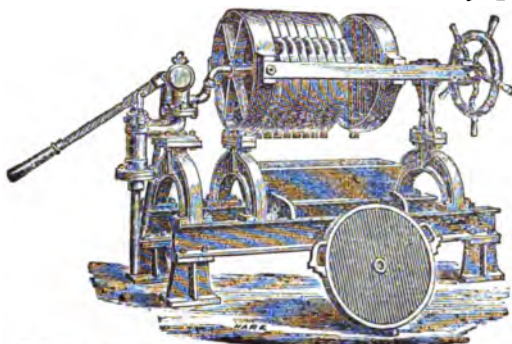
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THE CHEMICAL NEWS.

VOL. XXXIX, NO. 1005.

NOTE ON STEEL WELDING.

By SERGIUS KERN, M.E., St. Petersburg.

In some articles inserted in this journal the author stated that pure steel, nearly free from phosphorus and sulphur and containing 25 to 30 per cent of carbon, stands easily the process of welding, if, indeed, the work is done with care and by clever workmen.

It may be mentioned here that to a steel ship-plate, 2 ft. wide and $\frac{1}{2}$ inch thick, a steel plate ($2' \times 2' \times \frac{1}{8}''$) was easily welded, and a perfectly clean and good joint was received.

In another case steel strips ($6'' \times 4'' \times \frac{1}{8}''$) containing 25 to 26 per cent of carbon were welded together; very often after cooling the plate was bent double, through the weld, without the least fracture in or near the welded part. In some experiments such plates were bent at a dark heat, and they often, not always, resisted this severe test, as it is known that at this temperature the steel is more liable to break. These trials show that Russian Bessemer steel is of a very good quality.

TENACITY OF STARCH.

By GEORGE WHEWELL, F.I.C., F.C.S.

In consequence of the large quantity of farina (potato starch) which is used for the sizing and stiffening of yarn and cloth, the question often arises—Given several samples of starch, which will make the stiffest cloth and which has the greatest tenacity? To my mind the best method devised so far is the one used by Shier, and is as follows:—24 grains of the sample of starch are mixed with 400 grains of distilled water, and boiled with constant stirring for three minutes, then poured into conical test-glasses and allowed to stand for two hours; at the end of which time a flat metal disk seven-tenths of an inch in diameter, and not weighing less than 50 grains, is placed on the jelly and weights added until the skin is broken and the disk sinks.

In working the above method I find a difficulty in manipulating the weights on account of the smallness of the disk. If after a 50- or 100-grain weight has been placed on the disk you attempt to place a few smaller ones on also, the pressure is exerted unevenly, and the jelly is cut by the edge of the disk and not broken uniformly. I have devised the following modification, which I have used for the last three years, and find it to answer very well. The principle is to ascertain the number of grains required of any sample of starch which, when made into a jelly, will support a 100-grain weight for five minutes without breaking the skin. I have examined a large number of samples of farina, and find the number of grains required is from 18 to 28.

To determine the tenacity of a sample of farina I take, say, six conical shaped test-glasses 1 ounce capacity, and label the feet 18, 20, 22, 24, 26, and 28. Using a 2-ounce evaporating dish, I weigh out 18 grains of the farina, mix it with 26 c.c. of distilled water, and boil with constant stirring for three minutes. Pour it into the test-glass marked 18, and shake to make the surface of the jelly level, book the time, and allow to cool for exactly two hours. At the end of this time I take a 100-grain weight (mine being thirteen-twentieths of an inch in diameter), and place it on the jelly. If it is supported for more than

five minutes I place it on each of the others in succession until I come to the one through which it sinks, then I have got the tenacity to within 2 grains. The intermediate number will be found to support the weight for exactly the right time. The number of grains required I call the tenacity. If the tenacity of one sample is 18 and the other 28, it means that it requires 28 lbs. of one sample to give the same stiffness as 18 lbs. of the other.

NOTE ON CHLORIDE OF CALCIUM.

By O. GLUGE, Sarrebrück.

THE manufacture of soda by the ammonia process has, during the last few years, been much developed, owing to the advances made by Mr. E. Solvay, and is likely to become more generally employed. It is possible that it may ultimately supersede Leblanc's method, as improvements are being constantly introduced into it.

The reaction which occurs in this manufacture, as all your readers are aware, takes place between bicarbonate of ammonia and chloride of sodium. Bicarbonate of soda is precipitated and hydrochlorate of ammonia remains in solution. The bicarbonate is separated by filtration, and the solution is employed to furnish fresh bicarbonate of ammonia. This is done by distilling the solution with lime; then ammonia is disengaged and a strong ley of chloride of calcium is left in the distilling apparatus.

I am anxious to direct the attention of your readers to this latter product, which is formed in considerable quantity and which is commonly thrown away as useless, as it has hardly been employed hitherto in manufacturing processes.

It would be very desirable that men of science and manufacturers should endeavour to ascertain if a more extended use could be made of a product which can be got at such a cheap rate. The manufactures of soda by the process above mentioned can furnish chloride of calcium in quantity, in solution, in crystals, or even in the fused state to diminish the cost of carriage.

ON THE DETERMINATION OF SPECIFIC GRAVITIES.

By S. F. PECKHAM.

In the *CHEMICAL NEWS* (vol. xxxviii., p. 300) I notice an excellent article "On the Determination of Specific Gravities." The author's points are well taken in reference to the importance of some uniform method either of performing the operation or of confuting results, and his suggestions could with great propriety be referred to some committee of one of the Societies for a report and recommendations that might be generally adopted.

Having occasion some years ago to take the specific gravity of some small quantities of distillates from petroleum products, I was led to adopt the following method, which was found to be entirely successful with so small a quantity as 3 cubic centimetres.

I prepared a cube of aluminium, which was intended to be of 1 c.c., but was accidentally obtained smaller. This was suspended by a platinum wire of the smallest size, and weighing only a few milligrams. The wire was first weighed, and then attached to the cube, which was then weighed in air, and then weighed just immersed in distilled water, and then in the oil. The difference between the weight in the air and in water gave the weight of the water; the difference between the weight in air and in oil gave the weight of the oil. The weight of the oil divided by that of the water gave the specific gravity of the oil.

This method admits of perfect control of the volume of

the liquids compared, and of just as perfect control of the temperatures as any other method. Moreover, it admits of very rapid execution and of operating on very small quantities of liquid. For taking the specific gravity of all ordinary oils and other non-corrosive liquids a cube of aluminium would be superior to a cube of platinum, on account of its lower specific gravity, and its permanence in the air renders it superior to any other of the so-called base metals.

For the class of liquids named I have found the results obtained by this method more satisfactory than those obtained by the bottle when the quantity available was unlimited; I therefore recommend the method without hesitation as applicable to very small quantities of liquid, as reliable, and as incomparably more rapid of execution than any other method with which I am acquainted.

It has often been my custom when ascertaining specific gravities for purposes of comparison to cool the oils to zero Centigrade. This temperature is very easily obtained by immersing the vessel in fragments of ice placed in a funnel from which the melted ice-water can readily flow away. It may not be generally known that water in which pieces of melting ice are floating will rarely cool a vessel to zero, usually not below two degrees. If a tripod holding the funnel is placed over the balance-pan, and a funnel is selected with a long neck bent nearly at a right angle, the water from the melting ice may be discharged into a beaker to one side, while the vessel, holding only a few c.c. of liquid is cooled completely to zero with the least possible trouble.

For purposes of comparison this arrangement furnishes results often of great value in technical operations in which rapidity of execution is often of more importance than that fastidious regard for accuracy which cannot be overestimated in questions relating to the absolute "constants of nature."

Chemical Laboratory, University of Minnesota,
January 31, 1879.

ON THE NEW SUBSTITUTE FOR LITMUS.

By GREVILLE WILLIAMS, F.R.S.

WHILE engaged in preparing a Supplement to my "Hand-book of Chemical Manipulation" I made inquiries in several directions, with the view of ascertaining what improvements had recently been made in the volumetrical determination of acids and alkalies. Amongst the persons consulted was my friend Dr. Otto N. Witt, who kindly pointed out to me that the "orange 3" of Porrier formed an excellent substitute for litmus. This new coal-tar derivative was introduced into commerce about two years ago: I believe, however, that its manufacture has been almost entirely, if not quite, abandoned in favour of "orange 4" or tropæolin OO. This "orange 3" had, before its introduction into commerce, been studied independently by Dr. Witt and M. Griess; and M. Lunge, in the *Berichte der Deutschen Chemischen Gesellschaft*, expressly mentions it as an indicator. Its constitution will be apparent from its chemical name, dimethyl-amido-azobenzol-sulphonate of ammonium.

Lunge has shown that, under certain circumstances tropæolin and "orange 3" are superior to litmus, as they are unaffected by carbonic acid and sulphuretted hydrogen; they are therefore especially valuable in the analysis of soda residues.

As this matter possesses considerable interest and importance to those who have much to do with the determination of acids and alkalies, I have subjected "orange 3" and litmus to a careful comparison. I selected it instead of tropæolin, as it is much more delicate in its reactions.

Different specimens of litmus somewhat vary in their sensitiveness to acids and alkalies; that with which I

worked on this occasion ceased to indicate with sulphuric acid when more dilute than 1 to 50,000. The new indicator, on the other hand, remained sensitive to 1 part of sulphuric acid in 100,000 of water.

The solution of the indicator which I employ is made by dissolving 1 centigramme in 100 c.c. of distilled water, and contains therefore 1 part in 10,000. Even at this great dilution the liquid is of a full orange colour. It is worthy of remark that 1 centigramme of the substance is sufficient for five hundred determinations of acid or alkali; its cost is consequently inappreciable. It has also the advantage over litmus that its solution is less liable to decomposition on keeping.

The quantity of the solution of the indicator, of the strength given above, to be used in each determination is from one to two-tenths of a cubic centimetre. When this small quantity is added to a solution containing an alkali to be tested, there is no perceptible colouration as long as it is alkaline; but the faintest trace of acid turns the solution a pale but distinct pink tint, which is easily seen if the beaker be placed upon a sheet of white paper.

The best way of applying the test is, after each addition of acid from the burette, to allow a drop of the indicator to fall on the surface of the liquid in the beaker after the contents have been well stirred. By this mode of proceeding the indicator is distributed over a smaller space, and the reaction is therefore more distinct.

An immense advantage which this indicator possesses over litmus is that it is entirely unaffected by carbonic acid; the solution need not therefore be boiled, and the operation consequently takes much less time; in fact, four operations can be made in the same time as one with litmus. That without boiling the solution it competes perfectly with litmus as regards accuracy may be judged from the following table:—

Determination of Carbonate of Sodium Volumetrically.

One c.c. of the standard sulphuric acid = 0.051176 grm. of carbonate of sodium.

Carbonate of Sodium taken. Grms.	Standard Sulph. Acid used. Cub. centims.	Percentage of Carb. Sodium obtained.	Indicator used.
0.4655	9.07	99.71	Litmus.
0.9345	18.20	99.68	
1.0646	20.75	99.75	Orange 3.
0.6739	13.15	99.86	

In order to determine whether the new indicator gave as accurate results with ammonia as with carbonate of sodium, the following experiments were made upon a solution diluted to 5 per cent for titration:—

Determination of Ammonia Volumetrically.

One c.c. of the standard hydrochloric acid = 0.00697 grms. of ammonia.

Ammonia taken. Cub. centims.	Standard Hydrochloric Acid used. Cub. centims.	Percentage of Ammonia obtained.	Indicator used.
20	28.50	19.86	Litmus.
16	22.70	19.78	
18	25.50	19.75	
18	25.60	19.83	Orange 3.
14	19.85	19.77	
22	31.30	19.83	

It is obvious, therefore, that the new indicator can be employed in the determination of ammonia instead of litmus, and has many advantages over it.

The Banana.—M. Corenwinder.—The total sugar in this fruit when ripe and sound exceeds 20 per cent, three-fourths of which are crystallisable. When over-ripe the total sugar is reduced to from 16 to 14 per cent, whilst the non-crystallisable sugar rises to 11 or 12, thus being not merely relatively but positively increased.—*Comptus Rendus*.

EXAMINATION OF CAOUTCHOUC GOODS.

By DAVID LINDO.

It was stated some time back (Scientific Notes, *Quarterly Journal of Science*, April, 1877) that the use of oxide of zinc in the manufacture of caoutchouc nipples for milk-bottles has almost been abandoned.

This led to the examination of some caoutchouc nipples lately imported from England, and supplied by first-class houses. The examination was afterwards extended to samples of tubing supplied for chemists' use, and a few other caoutchouc goods.

Some specimens of chemists' tubing were found to consist solely of india-rubber and sulphur. Others contained a large proportion of carbonate of lime and a considerable quantity of oxide of zinc, also a little siliceous matter—apparently French chalk. Oxide of zinc was present in very large quantity in all the other goods, and siliceous matter never altogether absent.

The following were more particularly examined:—

- Lot No. 1.—Oval-shaped nipples, stamped F.
 " No. 2.—Oval-shaped nipples, stamped J.
 " No. 3.—Cylindrical nipples, numbered by maker 2, 3, 4.
 " No. 4.—Ring for teething infant.
 " No. 5.—Caoutchouc doll.
 " No. 6.—Tubing for chemists' use, $\frac{1}{2}$ in. diameter.
 " No. 7.—" " " $\frac{1}{4}$ in. "

The ash was estimated in all these samples, and the oxide of zinc. The quantity of carbonate of lime in Lots 5, 6, and 7 was also ascertained.

The zinc estimates were made from the material direct by cutting up the latter very small, and adding one piece at a time to a mixture of pure carbonate of soda and nitre kept in fusion in a porcelain crucible. The mass was dissolved out with moderately dilute acetic acid, the zinc precipitated with sulphuretted hydrogen, the sulphide zinc converted into chloride; precipitated with carbonate of soda, and weighed as oxide.

When two estimates of the same sample were made, different weights of material were employed.

Lot No. 1.

External surface smooth and not easily abraded. Composition not quite uniform. Ash, mean of three experiments, 46.79 per cent. Oxide of zinc from average sample, 44.14 per cent.

Lot No. 2.

Surface smooth and not easily abraded. Composition uniform. Ash, 47.22 per cent. Oxide of zinc, two estimates:—

1st.	43.52 per cent.
2nd.	43.68 per cent.

These nipples contained a little more clay than Lot No. 1.

Lot No. 3.

These nipples exhibited considerable difference in percentage of mineral matter. Their surfaces were quite smooth, but could be abraded rather easily with the finger-nail.

Makers' Numbers.	Ash, Per cent.	Oxide Zinc, Per cent.
2	62.70	60.71
3	52.58	48.89
4	51.68	49.63

Lot No. 4.

Surface very rough and easily abraded. Ash, 32.07 per cent. Oxide of zinc, two estimates:—

1st.	29.01 per cent.
2nd.	29.18 per cent.

Lot No. 5.

Surface very rough and easily abraded. Composition not quite homogeneous. Ash, estimated from sample cut

from different parts, 65.58 per cent. Carbonate of lime, 11.10 per cent. Oxide of zinc, two estimates from average sample:—

1st.	49.98 per cent.
2nd.	49.90 per cent.

Lot No. 6.

Ash, 49.12 per cent. Oxide of zinc, 15.47 per cent. Carbonate of lime, three estimates:—

1st.	30.21 per cent.
2nd.	30.22 per cent.
3rd.	30.45 per cent.

Lot No. 7.

Ash, 48.90 per cent. Two estimates of carbonate of lime and oxide of zinc:—

	Carbonate of Lime, Per cent.	Oxide of Zinc, Per cent.
1st.	30.50	15.34
2nd.	30.82	15.33

Certain acids, when brought in contact with these compositions, were found to produce some curious phenomena, which will be fully described further on.

The acids employed were—

Sulphuric acid	sp. gr. 1.828
Hydrochloric acid	" 1.154
Nitric acid	" 1.374
Acetic acid, containing 63 to 64 per cent anhydrous acid.	

They were employed of the above strengths, and also much weaker, the dilutions being carried out by volume.

The ounce will be understood to mean a fluid ounce. The phenomena alluded to are—

1. Distension of the material from absorption of fluid.
2. The appearance of small elevations on the surface.
3. Crimping of the edges.

These effects appear to be due to the following causes:—

1. The acid has a strong tendency to enter the pores of the material, in consequence of its affinity for the lime or oxide of zinc.
2. Certain acids (notably acetic acid) not only acts solvent on the mineral matters, but also softens the caoutchouc, rendering it in consequence more easy of distension, and at the same time diminishing its power of contracting after distension. The minute state of division in which the vulcanised rubber is left after disintegration of the mineral matter seems to favour this action.
3. The acid entering the pores of the softened material more readily than the solution of zinc or lime salt formed within the pores escapes from them, distension follows.
4. Small elevations will generally appear on the surface of the material, because the distension at first is merely superficial. These elevations are also sometimes produced by escaping gas-bubbles.

If a sharp edge is presented to the action of the acid, crimping of the edge will result, and remain apparent until the action has proceeded to some depth below the surface of the material, and distension has in consequence become more general.

The general effects produced by the acids are described below, and a few examples given of the absorption occasioned by acetic acid, &c. Strictly comparable experiments could not be made with the samples at my disposal (except perhaps with the chemists' tubing), on account of the irregular form, unequal thickness, and various degrees of elasticity.

The appearance which most samples present under the action of acetic acid is very beautiful, especially if observed with a lens. The phenomenon soon becomes apparent to the naked eye, except with extremely dilute acid; but a lens is required to observe it in the early stage, and to

render visible that fine crimping of the edges which constitutes the only apparent effect produced by very weak acid. With the aid of a good watchmaker's eye-glass I have never failed (in numerous trials) to detect the presence of 1 part of anhydrous acetic acid in 3000 parts of water, by immersing a small piece of nipple (weighing 2 or 3 grains) from Lot No. 1 in 10 c.c. of the diluted acid. Distinct crimping was observed with the lens after twelve to twenty-four hours, when the experiment was performed at the ordinary temperature of the laboratory* (78° to 82° F.). If the fluid and sample were heated on the water-bath, in a closed bottle, the reaction was generally obtained in two or three hours.

It is probable that formic acid will be found to produce the same effect, and that the reaction may be rendered available as a means of detecting minute quantities of either acid in the free state in the absence of the other, provided free mineral acids in large quantity are also absent.

I should here observe that all samples of the material will not be found equally sensitive to the action of the acid, a marked difference being observed in some instances between two samples of apparently the same composition.

Action of Sulphuric Acid.

The strong acid decomposes the material in a few hours. Some specimens were observed to crimp slightly when first immersed, others not at all.

Acid 1 in 2† was found to dissolve out oxide of zinc pretty freely, except from those samples which contained a notable quantity of carbonate of lime. After twelve days' immersion in acid of this strength the elasticity of the material seemed but slightly impaired. Not the slightest crimping was produced, nor was weaker acid found to occasion it. Acid 1 in 20 dissolved out enough zinc from most samples to be readily detected in the fluid after thirty-six hours' immersion.

Action of Hydrochloric Acid.

A piece of No. 1 was immersed in about four times its weight of strong acid. In half an hour a notable quantity of zinc was found in solution. Pieces of No. 1 and No. 5 were immersed in excess of strong acid and examined with lens. Numerous gas-bubbles from No. 5, and a few from No. 1. Cut edges acquired a reddish tinge. After two days both samples had acquired a reddish colour all over. Decided crimping. Sulphur had separated. Some distension of material generally. After four days removed samples from acid, and washed. Elasticity somewhat impaired. On cutting, it was seen action had advanced very far into the material. Original thickness about $\frac{1}{4}$ th inch. A nipple from No. 1 was cut in two lengthwise, and immersed in 2 ozs. acid, 1 in 5.

After nine days faint traces of crimping observed with lens. After fourteen days these had nearly disappeared. Removed sample. Washed, dried with towel, and weighed.

Weight	25.17 grains.
Original weight	24.19 "
	0.98 gr.

Adhering moisture would account for the difference. Quantity of oxide of zinc found in *one* ounce of the fluid, 0.22 gr. Material seemed uninjured.

Acid 1 in 20 failed to produce the least appearance of crimping with the most sensitive samples, the specimens being kept under the action of the acid for a month, and examined carefully with the lens almost daily.

* The temperature at which all the experiments were performed except when otherwise stated.

† One volume of sulphuric acid, sp. gr. as stated, mixed with an equal volume of water. The same for all degrees of dilution, and for all the other acids employed.

Action of Nitric Acid.

The strong acid reduces the material to pulp in a few hours. Acid 1 in 5 also effects decomposition, though much more slowly. A piece of No. 1 weighing 20 grains was immersed in half an ounce of acid of this strength, and finally examined after eighty hours (slight crimping had been observed in the interval). On cutting, decomposition was found to have advanced to a considerable depth, leaving apparently only sulphur on both surfaces, which formed a brittle crust. Acid 1 in 10 appeared to have little action beyond dissolving out the metallic oxides and slightly softening the caoutchouc, producing crimping and general distension of the material. No effect of this kind could be obtained with acid weaker than 1 in 100, and very careful inspection with a strong lens was required to observe the faint traces of crimping produced when the most sensitive samples were immersed in acid of this degree of dilution.

Action of Acetic Acid.

Acid of the strength employed in these experiments (or weaker) was found to have very little action on pure vulcanised rubber. Pieces of tubing (composed of the latter) left floating in the acid for days appeared quite unaltered. On mixing the fluid with water, however, it produced milkiness, showing that a small quantity of caoutchouc had gone into solution. The fact that acetic acid has very little action on pure vulcanised rubber renders its extremely energetic action on the mineralised article truly remarkable, especially when it is remembered that as compared with mineral acids its affinity for bases is feeble.

The strong acid dissolves out the mineral matter very readily, and produces distension so rapidly that no elevations appear on the surface of the material, and very slight crimping of the edges is observed.

Distension, however, although it occurs more rapidly with strong than with weak acid, is often sooner arrested with the former than with the latter, and long before the acid has become exhausted. It appears to cease as soon as the fluid contains a certain proportion of salts in solution. From this it follows that a weak acid, other things being equal, may produce as great or greater distension after a time than a strong one. There are, however, so many circumstances that might influence the result (notably the degree of elasticity possessed by the material) that I should hesitate to accept this as the correct view without further and more exact experiments than I have as yet been able to undertake.

The following experiments serve principally to illustrate the bare fact that very considerable distension does take place with the acid, whether weak or strong:—

A piece of nipple from Lot No. 2, weighing 28.72 grs., was immersed in 5 ounces of acid, 1 in 300. After a month the sample was removed, and placed in 5 ounces of fresh acid of the same degree of dilution. The sample was weighed forty-six days after the commencement of the experiment.

Weight, 112.18 grs. Apparent gain, 83.46 grs.

The material could be torn rather easily. A portion ignited left a considerable residue of oxide of zinc.

Lot No. 2, piece weighing 10.83 grs. Lot No. 6, piece weighing 8.10 grs. Immersed together in 1 ounce of the strong acid. Weights after twenty-one days:—

No. 2, 19.80 grs. Apparent gain, 8.97 grs.
No. 6, 17.30 grs. " " 9.20 grs.

Lot No. 1.—Nipple cut in two lengthwise. Weight, 23.59 grs., immersed in 2 ozs. acid, 1 in 10. Weight after fourteen days:—

93.60 grs. Apparent gain, 70.01 grs.

Quantity of oxide of zinc found in 1 ounce of the fluid, 3.78 grs. Material very tender.

Lot No. 1.—Nipple cut in two lengthwise. Weight, 27·89 grs. Immersed in 2 ozs. acid, 1 in 50. Weight after fourteen days :—

138·70 grs. Apparent gain, 110·81 grs.

Quantity of oxide of zinc found in 1 ounce of the fluid, 1·43 grs. Material very tender.

Lot No. 2.—Nipple cut in two lengthwise. Half immersed in 2 ozs. acid, 1 in 10; the other half in 2 ozs. acid, 1 in 50. Original weight of half in acid 1 in 10, 19·97 grs. Weight after fifteen days :—

73·30 grs. Apparent gain, 53·33 grs.

Original weight of half in acid 1 in 50, 15·19 grs. Weight after fifteen days :—

72·80 grs. Apparent gain, 57·61 grs.

The halves presented originally very nearly, if not quite, the same extent of surface, the difference in weight being due to difference in thickness. After being acted on by the acid, the material in both experiments was found extremely tender.

After this considerable distension fluid escapes so rapidly from the pores of the material, on removal from the acid, that the weight obtained only approximately represents the amount of absorption that has taken place.

The amount of mineral matter dissolved out must also be taken into account. If left to dry these distended samples gradually contract to about their former dimensions, but never recover their elasticity unless the action of the acid has been only slight.

In reference to the phenomena of crimping, &c., by far the finest effects are produced by a dilute acid, say 1 in 300. The result obtained in the following experiment (one out of a very large number tried) will serve to illustrate the action of acid of this strength on a fairly sensitive sample of the material :—

Lot No. 1.—Nipple with top cut off; weight 17·9 grs.

Lot No. 6.—Piece of tubing cut off; weight 9 grs.

Immersed both in $\frac{1}{2}$ oz. acid, 1 in 300. Examined after twelve hours :—

No. 1.—Decidedly but finely crimped on cut edge, quite apparent to naked eye. Surfaces slightly raised. A few bubbles.

No. 6.—Edges slightly crimped. Requires lens to see distinctly. Surfaces very slightly raised.

After thirty-six hours :—

No. 1.—Beautifully and finely crimped. Base ring looks like fine sponge. Exterior surface covered with small round elevations, beautifully regular. Interior surface very slightly raised. No bubbles.

No. 6.—Edges rather strongly crimped. Surface slightly raised. No bubbles.

Very fine effects can be obtained with much weaker acid than 1 in 300, provided the material is sufficiently sensitive. With most of the nipples I have obtained reaction in acid 1 in 2000 (equal to about 1 part anhydrous acid in 3000 parts of water). With sample Lot No. 5 acid 1 in 300 hardly produced any effect.

Old samples that have lost their elasticity to a considerable extent will be found equally inactive.

Falmouth, Jamaica,
January 22, 1879.

THE ELECTRIC LIGHT IN THE BRITISH MUSEUM READING-ROOM.

LAST Monday will henceforth be looked upon as an interesting point of departure in the history of the British Museum Library, for on that day its manifold treasures were for the first time thrown open by night as well as by day to those entitled to use them.

For the past three weeks Mr. Bond, the Principal Librarian, and M. Berly, C.E., the London representative

of the Paris Société d'Electricité, assisted by their respective staffs, have been making repeated experiments on the practicability of lighting up the British Museum Reading-Room by means of the Jablochkoff system of electrical illumination. Having partially determined upon the number and position of the lamps to be used, Mr. Bond decided that on the Monday, Tuesday, and Wednesday of the present week the Reading-room should be kept open until seven o'clock, so that the holders of reading-tickets might have the opportunity of practically testing the value of the welcome innovation. On Monday evening, accordingly, about two hundred readers remained behind after the usual hour for closing, and when, at a few minutes before six o'clock, the twelve Jablochkoff candles in shades of opal glass suddenly burst into light, those present forgot for the moment that they were in a building devoted to silence and study, and evinced their approval of the efforts of the Museum officials on their behalf by breaking into a burst of applause, a sound which we will venture to say has never before been heard beneath Sir Antonio Panizzi's famous dome.

Roughly speaking the Reading-Room is a circle, nineteen-twentieths of which are devoted to the public, the remaining twentieth forming the passage into the Library. In the centre there are three circular desks, the inner one being used for the delivery and return of books, and the two others, which are breast high, for stacking and using the voluminous catalogues. From these run radially nineteen desks divided lengthways by a partition, and lettered from A to T both inclusive, but missing Q, seventeen of which are double, the two end ones being single. At present the first four, A, B, C, and D, are each illuminated by a Jablochkoff lamp, placed on a standard fifteen feet high, fixed exactly in the middle of each desk, being sustained by the longitudinal partition which separates the readers, the remaining fifteen desks being lighted by seven lamps placed alternately. The remaining lamp is placed in the centre of the room, and lights the desks of the Superintendent and his assistants. The general opinion amongst the readers appears to be one of unanimous approbation of this mode of lighting. We have thoroughly tested the matter in a practical manner by reading, writing, tracing, drawing, and painting at one of the first four desks as well as at those which are only lighted alternately. In the first case there is abundant light for comfortable working at any part of the four desks, but in the latter a reader sitting at either end of the illuminated desks has to twist himself round most uncomfortably to get out of his own shadow. We venture to think, therefore, that for the new mode of lighting to be thoroughly satisfactory to all, the whole of the nineteen desks must each be provided with a lamp, thus rendering the imitation of daylight as perfect as need be.

It is agreed on all hands that the light is mellow and soft, and most agreeable to work by. Now and then, it is true, there is a sudden flutter in the light, and occasionally it waxes and wanes slightly, but these defects will no doubt disappear when everything is in full working order.

The source of electricity is a 20-light duplex Gramme machine of the latest construction, worked by a Robey portable engine of 16 horse-power nominal. There are four circuits of five lamps, but only sixteen are used at present; that is to say, twelve in the Reading-Room, one in the Entrance Hall, one under the portico, and two in the machine and engine-shed. The machine and its engine are placed outside in a wooden erection at the north-west corner of the Museum buildings, about 200 yards distant from the Reading-Room.

The four candles used in the lamps at desks M, O, R, and T are of an improved kind lately invented by one of M. Berly's assistants, and are now tried for the first time. They differ from the ordinary Jablochkoff candles in the insulating material between the carbons being replaced by a composition which we suppose must be a feeble conductor. Extinction, except for a moment, is therefore impossible. The use of the carbon bridge for

lighting is consequently entirely obviated. Not only this, but one, two, three, or the whole of the four candles may be lighted or extinguished at will simply by turning the handle of the commutator, or if one goes out it re-lights itself automatically without extinguishing its neighbours.

It would, of course, be premature to speak of the cost of permanently carrying out this immense boon to students and literary men generally; we may, however, mention that a reduction has recently been made in the price of the ordinary ninety-minute candles of something like 40 per cent. That the innovation is already highly appreciated is shown by the large attendance of real workers on the three evenings in question, and by the almost unanimous chorus of approbation indulged in by readers of all classes. The Société d'Électricité deserves great credit for the public spirit they have shown in gratuitously supplying everything necessary for making this interesting experiment.

Mr. Bond and his able coadjutors seem determined to extend the use of the treasures under their charge in every possible direction, and it ought to be the duty as well as the pleasure of the literary, artistic, and scientific press of this country to strengthen their hands by generously commending and seconding their well-intentioned efforts.

PROCEEDINGS OF SOCIETIES.

PHILOSOPHICAL SOCIETY OF GLASGOW.

CHEMICAL SECTION.

Ordinary Meeting, February 10, 1879.

Mr. J. J. COLEMAN in the Chair.

THE first paper was read by Dr. JOHN CLARK, "*On the Action of Phosphuretted Hydrogen on the Animal Organism.*" His experiments were carried out in conjunction with Dr. Henderson. Small animals, usually rats, were introduced into the jar. There appeared to be considerable difficulty in breathing, and itchiness of the skin. Death took place usually within half an hour. A very small amount of gas has a fatal action; even 1 part in 5000 causes death. The blood was dark and venous, and the lungs inflamed. Portions of the liver and blood were tested for phosphorus compounds by introducing them into a flask in which hydrogen was being evolved from zinc and hydrochloric acid. The flame of the hydrogen had a green colour, and exhibited a spectrum of green bands, characteristic of phosphorus.

Mr. TATLOCK read the second paper, "*On Magnetic Iron Sand from the Kyles of Bute.*" This sand forms black streaks on the Argyllshire shore. It is separable into two portions by the magnet, both of which have the same appearance and crystalline form; one is magnetic, and the other not. On analysis the magnetic portion was found to contain 83.55 per cent of Fe_3O_4 and 15.6 per cent of Fe_2O_3 ; and the non-magnetic portion contained 0.67 per cent of Fe_3O_4 , and 97.64 per cent of Fe_2O_3 . If the magnetic variety had been oxidised to the non-magnetic variety no disintegration of the crystal had taken place.

The last paper was read by Mr. J. B. HANNAY. The subject was, "*Variation in the Magnetic Constituents of Minerals.*" It has been observed that many rocks are feebly magnetic. The author's method of determining the amount of this magnetic attraction has been elsewhere described. It consists of suspending a sample of the rock to the beam of a balance, and determining the amount of oscillation produced by a magnet placed above the pan. The magnetic constituents of these rocks, comprising specimens of pyrochlor-obsidian, natrolite, rhodonite, and gneiss, were separated by pulverisation and

attraction by a magnet. The portion separated was again reduced to powder, and the magnetic portion again separated, and so on till no non-magnetic residue remained. The magnetic portions all had almost the same specific gravity, and after deduction of a silicate of ferroso-ferric oxide, $\text{Fe}_3\text{O}_4 \cdot 2\text{SiO}_2$, the residue was found to approach the composition of a spinel, composed of an oxide of the formula $\text{M}_2\text{O}_3 \cdot \text{MO}$. This residue was magnetic, when iron constituted either one or other of the oxides, the remaining oxide consisting of alumina or lime. Mr. Hannay's conclusion, therefore, is that such a mixed oxide, or spinel, may possess magnetic properties when associated with magnetic silicate of ferroso-ferric oxide.

EDINBURGH UNIVERSITY CHEMICAL SOCIETY

February 12, 1879.

Mr. GEORGE MACGOWAN, F.R.S.E., in the Chair.

A PAPER was read by Dr. W. INGLIS CLARK "*On the Action of Chlorinated Substances on Alcohol,*" in which he described the practical method of preparing chloroform, and the chemical reactions which most probably occur in its formation.

CORRESPONDENCE.

THE INSTITUTE OF CHEMISTRY.

To the Editor of the Chemical News.

SIR,—When the Institute of Chemistry was first proposed the promoters gave great offence by holding hole-and-corner meetings of their personal friends, instead of inserting an advertisement in your columns asking all professional chemists to meet together and discuss their mutual wants with a view of forming an association.

Whatever excuse there might have been at that time for such a course of procedure, it is surely unnecessary and undesirable to continue the same system.

Throughout the year 1878 the Members of the Institute heard of absolutely nothing being done by the Council to advance the interests of the profession, until in December—just before the new subscriptions fell due—the Institute showed a faint sign of life, and a meeting was held, at which the important subject of "Trade Certificates" was discussed.

This seemed a step in the right direction, and we looked forward anxiously for a verbatim report of the proceedings. With a large number of Members scattered all over the kingdom, and with ample funds for the purpose, our Council surely could not have contemplated anything less.

Judge then of the disgust of myself and other subscribers when it leaked out that the executive dared not publish a report of the meeting because nearly all the speakers upheld the system of giving certificates for advertising purposes. They were expected to curse, but verily they blessed them altogether. But why should the discussion not be published? Have the executive discovered that they have been cherishing a snake? Are some of the worst sinners to be numbered amongst the very elect, i.e., the Council of the Institute of Chemistry of Great Britain and Ireland?

Now, Sir, I protest against this hole-and-corner business. Every Fellow of the Institute has a *right* to know all that occurs at the public meetings of his society, and the attempt to prevent publicity of discreditable proceedings will, if persisted in, speedily prove the ruin of the Institute.

In a few days' time a second conference will be held, at which the subject of the "Adulteration of Food and

Drugs" will be discussed. While deprecating the choice of Dr. Voelcker as the leader of the discussion—for, eminent as he is in his own department, his opinions command no respect among those chemists who have had special experience in the examination of food and drugs—it is to be hoped that his speech and that of all others who take part in the debate will be fully reported and circulated among the Members.

As the Council of the Institute allowed the year 1878 to pass without moving a finger to advance the interests of the profession, and as there are ample funds in hand, I had hoped they would have had the discretion not to make any call for 1879. As it is, I shall wait to see what the "Government Programme" is before I pay my subscription, and I hope the other Members will do the same. In short, I challenge the Council to show what they intend to do with the money now in hand.—I am, &c.,

A DISGUSTED PROMOTER.

THE BESSEMER PROCESS FOR TREATING SULPHIDES.

To the Editor of the Chemical News.

SIR,—With reference to the remarks on my experiments in the CHEMICAL NEWS, vol. xxxix., p. 94, that "The analyses made seem to show that in the regulus obtained there is often much less sulphur than corresponds to the formula FeS —a fact that points to a lower sulphide of iron than any hitherto recognised," Is it not possible the excess of iron present may be accounted for in another way, viz., by the presence of metallic iron?

I propose making the following experiment in a Bessemer provided with a lime lining, viz.—To Bessemerise FeS with a hot blast and without adding silica, and I expect (provided the temperature is sufficiently high) that the protoxide of iron formed will react on the protosulphide of iron and produce metallic iron with the evolution of sulphurous acid.—I am, &c.,

JOHN HOLLWAY.

7, Jeffrey's Square, St. Mary Axe, London, E.C.,
March 1, 1878.

ALUM IN FLOUR AND BREAD.

To the Editor of the Chemical News.

SIR,—Mr. Penney, in his article on "Alum in Flour and Bread" (CHEMICAL NEWS, vol. xxxix., p. 80) remarks that the logwood test for alum is to be thoroughly relied on as a qualitative test if properly applied, and that it is of little consequence whether the solution is old or new. In this conclusion I quite agree with him, and I wish to point out a reason for its sometimes failing.

I have noticed that if the mixture of logwood and carbonate of ammonia solutions is allowed to stand a short time before applying it to the flour or bread containing alum, it imparts a dirty brown colour, and sometimes makes no change in the colour, according to the length of time it has stood, but if applied immediately it never fails to give the blue colour.—I am, &c.,

ALFRED J. M. EDGER.

Laboratory, 13, Dean Street, Newcastle-on-Tyne,
February 27, 1879.

THE DISCUSSION ON WATER ANALYSIS AT THE CHEMICAL SOCIETY.

To the Editor of the Chemical News.

SIR,—Will you allow me to repeat what I said at a recent meeting of the Chemical Society, since, from my words not being fully reported, an inaccurate statement was attributed to me, and, furthermore, because I should like

to add one or two remarks which I omitted on the occasion referred to.

I drew attention to the fact that Prof. Dittmar had very carefully described the details of a method of burning water residues in the CHEMICAL NEWS about a year ago. I stated that, having tried this method upon some ten or twelve waters, I had found it yield very accurate results. These waters contained exceedingly small quantities of any solid constituents, and especially little organic matters. I described then the process as I had used it, saying that the nitrogen was burned with soda-lime in a current of hydrogen. Will you allow me to add that the ammonia, instead of being collected in acid, was absorbed by pure distilled water, and with perfect condensation. Prof. Dittmar mentions the constant presence of ammonia in mineral acids, and I experienced a serious difficulty in attempts to prepare acids which, after neutralisation with pure sodic carbonate, would not give a strong colouration with the Nessler test.

The soda-lime used, both to absorb the carbon dioxide and to burn the organic matter for nitrogen, was prepared from pure soda and ignited marble. It was finely pounded, and passed through one sieve on to a finer one, so as to free the material from dust, and at the same time remove all pieces larger than a pin's head.

I believe this combustion process will satisfy the requirements of all those who have only an occasional need of such a method of analysis.—I am, &c.,

W. N. HARTLEY.

King's College, March 4, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 5, February 3, 1879.

Remarks on the Third Reply of M. Pasteur.—M. Berthelot.—A continuation of a stale and unprofitable discussion.

Fermentation of Cellulose.—P. Van Tieghem.—It does not seem that there is a diastase of cellulose formed in excess by amylo-bacteria and acting at a distance from it. Microscopic observations show that the solvent action of the amylo-bacteria upon cellulose is produced by direct contact. If the hypothesis of a diastase naturally offers itself to the mind to explain the first stage of the fermentation of cellulose and in general of the insoluble matters produced by living beings it must be recognised as not readily verifiable.

Liquefaction of Silicide of Hydrogen.—M. Ogier.—The author has performed this experiment with the apparatus of Cailletet. At ordinary temperatures (about 10^5) liquefaction does not take place under pressures of 200 to 300 atmospheres. On the contrary, from 50 atmospheres the cooling due to the release determines the production of a thick mist and of a manifest trickling of liquid down the sides of the tube. Under these conditions the gas is at a temperature bordering upon its critical point. It suffices, indeed, to cool it a few degrees below zero in order to effect a total condensation. Hydric silicide is liquid at -11° at the pressure of 50 atmospheres; at -5° under 70 atmospheres; at -1° under 100 atmospheres, whilst at 0° it remained liquid up to 200 atmospheres. The conditions of the liquefaction of this gas resemble those of marsh-gas, with which it has so many analogies.

Determination of Methyl Alcohol in Commercial Methylens.—C. Bardy and L. Bordet.—The authors point out the imperfections of the ordinary process founded on the transformation of methyl alcohol into methyl

iodide. They effect the conversion in a special apparatus, a figure of which is inserted in their memoir.

The Wagnerite of Bamle in Norway, and on a Retinite from Russia.—F. Pisani.—Wagnerite, a fluoriferous phosphate of magnesia originally met with at Werfen, in Salzburg, is identical with a mineral subsequently discovered at Bamle, in Norway, and provisionally named Kjerulfine. The latter, however, contains a percentage of lime. The Russian retinite was formerly mistaken for a manganiferous garnet.

No. 6, February 10, 1879.

On Fermentations, a Final Reply to M. Pasteur.—M. Trécul.

Fourth Reply to M. Berthelot.—M. Pasteur.

Hydro-electricity and Hydro-magnetism: Experimental Results.—C. A. Bjerknes.—The author has examined the action between two pulsating bodies; between a pulsating body and another which oscillates; and between two oscillating spheres. In all these cases concordant pulsations produce attractions, and opposed pulsations give rise to repulsion.

The Green Phosphorescent Light of the Molecular Shock.—W. Crookes, presented by Th. du Moncel.—An abstract of a paper recently read before the Royal Society.

Dissociation of Chloral Hydrate.—MM. Engel and Moitessier.—The authors, along with M. Wurtz, and in opposition to M. Troost, maintain that chloral hydrate does not exist as a definite gaseous compound, and that its equivalent does not correspond to 8 vols. The dissociation is effected in an atmosphere of chloroform at 61°.

Researches on Beer-yeast.—MM. Schützenberger and Destrem.—Yeast digested alone at 30° for twenty-four hours lost 1·77 per cent of solid matters. If digested along with sugar there is an increase of solids amounting to 11·3 per cent on the weight of the yeast.

Homologues of Oxyheptic Acid.—E. Demargay.—The author has prepared the following homologues:—oxytetric, oxyptenic, oxyhexic, and iso-oxyhexic acids.

Analysis of an Ethiopian Honey.—A. Villiers.—This honey, known by the natives as *tasma*, is found in subterranean cavities in Ethiopia, and is said to be collected by a species of mosquito. Its composition is:—

Water	25·5
Levulose and glucose free from)	
saccharose	32·0
Mannite	3·0
Dextrine	27·9
Ash	2·5
Sundries and loss	9·1

100·0

Process for Enriching Phosphates with a Carbonated Gangue.—L. L'Hôte.—The author heats to bright redness so as to expel the carbonic acid, and then removes the caustic lime formed by lixiviation with very dilute hydrochloric acid. [To the best of our belief a substantially identical process has been for many years in occasional use at the Great Eastern Chemical Works, Stowmarket.]

Bulletin de la Société Chimique de Paris,

No. 1, January 5, 1879.

Examination of some Mineral Waters of Auvergne.—E. Willm.—Not susceptible of useful abstraction.

Transformation of Starch into Glucose by Cold Water.—J. Riban.—The author had prepared, according to Mohr's directions, a solution of salted starch by boiling 1 part of powdered starch in 100 parts of water, saturating with common salt and filtering. After a year the solution became less and less sensitive to iodine, and after three or four years ceased to react with it at all. The liquid

remained limpid and neutral, and on microscopic examination showed no trace of any organised ferments, but it readily reduced cupropotassic reagent and turned yellow on boiling with acids. A special experiment with potassic ferricyanide showed that this reduction was due only in a small part to dextrin.

Determination of the Specific Gravity and Coefficients of Expansion of Liquid Methyl Chloride.—MM. Camille Vincent and Delachanal.—Noticed in the *Comptes Rendus*.

Remarks on the Memoir of E. and O. Fischer on the Constitution of the Rosanilins.—A. Rosenstiehl.—The author considers it as established that there exist no rosanilins containing in their molecule more than 20 atoms of carbon, and that there are three distinct rosanilins, viz., rosanilin- α , decomposable into aniline and paratoluydin, which MM. Fischer designate as pararosanolin, and prove that a single molecule of toluydin enters into its composition; rosanilin- β , decomposable into aniline and orthotoluydin; the authors might call it orthorosanolin; they have not studied, it but confound it with the following. Rosanilin- $\alpha\beta$ (Hofmann's rosanilin), resolvable into aniline, orthotoluydin, and paratoluydin. M. Rosenstiehl at the outset of his researches considered this last compound as a mixture of the two former, but subsequently was led to regard it as a well-defined chemical individual. The beautiful researches of E. and O. Fischer have removed all doubt on this point, and shown that it is the superior homologue of pararosanolin, and not its isomer. The second must be the isomer either of the first or the third.

No. 2, January 20, 1879.

Preparation and Properties of Potassic Cobaltocyanide and of Certain Derivatives.—A. Descamps.—Already noticed.

Memoir on the Curves of Solubility of Salicylic and Benzoic Acids.—E. Bourgoïn.—The curve of solubility of benzoic acid in water is analogous to that of salicylic acid. As far as about 35° the two curves are represented by parabolas closely approximating, but above this temperature they change their nature, becoming modified in the same manner. Benzoic acid, which was at first rather more soluble than salicylic acid, becomes less soluble, so that the two curves intersect each other at 40°, at which point the solubilities are of course exactly the same.

On Laurent's Carminaphtha.—A. Guyard.—The author finds that this compound, whose existence has been doubted, may be prepared by dissolving at a gentle heat 1 equivalent or 128 grms. of naphthalin in a sufficient quantity of glacial acetic acid. On the other hand, 12 equivs. or 600 grms. of chromic acid are also dissolved in a sufficient quantity of cold glacial acetic acid. He then adds at a gentle heat the chromic solution to the naphthalin until the mixture takes a green tint, then, when all the chromic acid has been introduced, he boils for a few minutes. If the whole is then saturated with alkali, or an alkaline carbonate, and the liquid acidulated afresh, the carminaphtha is precipitated in red or brown-red flocks. He concludes that carminaphtha is really a product of the oxidation of naphthalin by chromic acid. When it is formed little carbonic acid is evolved, whilst when phthalic acid is produced carbonic acid is evolved in quantity. Carminaphtha is a very stable compound and dyes wool and silk a deep reddish brown without mordant, the shades thus obtained being less remarkable for beauty than for permanence.

Russian Chemical Society: Session of Nov. 2/24, 1878.—M. Wagner presented researches by MM. Schirokoff and Saytzeff on allyl-diethyl-carbinol, and a communication by M. Sorokine on the oxidation of diallyl and on the hexylic glycol thus derived. M. Barsilovsky sent a memoir on the azo-derivatives of toluen. A paper of M. Ponomareff's was read, on compounds belonging to the

uric acid group. M. Menschoutkine laid before the Society the results of researches on the formation of compound ethers corresponding to the non-saturated tertiary alcohols. M. Gustavson communicated a paper on the addition-products formed by chloride of aluminium with benzine and toluene. M. Saytzeff gave a preliminary account of a hydrocarbon obtained by acting with dilute sulphuric acid upon allyl-dimethyl-carbinol.

Justus Liebig's Annalen der Chemie,
Band 194, Heft 2 and 3.

On Daubréelite, a New Meteoric Mineral.—J. Lawrence Smith.—This mineral, in a state of purity, consists of shining black fragments of more or less foliaceous structure, somewhat resembling molybdenite. It is not in the slightest degree attacked by hydrochloric acid, cold or hot, but dissolves slowly in warm nitric acid, without depositing sulphur. Its specific gravity = 5.01. Its composition is:—

Sulphur	42.69
Chromium	35.91
Iron	20.10

98.70

Researches on Ketons of the Aromatic Series.—W. Stadel.—This paper also is not adapted for abstraction.

Band 195, Heft 1 and 2.

Nitrosalicylic Acids and the Isomerisms of the Benzol Derivatives.—H. Hubner.—A very extensive memoir, utterly incapable of useful abstraction.

Researches on the Non-saturated Acids (Second Treatise).—Rudolph Fittig.—This memoir, which extends to upwards of 100 pages, comprises:—Further contributions to the knowledge of fumaric and maleic acids, by C. Petri; investigations on the oil of Roman camomile, which, again, consists of a paper, by H. Kopp, on the organic acids obtained on the saponification of the above-mentioned oil; one by Julius Köbig on the several ingredients of Roman camomile oil; one by A. Pagenstecher on angelic and tiglic acids; and a note by R. Fittig himself on the constitution of these two acids. The third grand division of the memoir is devoted to the non-saturated aromatic acids. Under this head R. Fittig and F. Binder describe the addition products of cinnamic acid; E. Posen treats of amido-hydro-cinnamic acid, otherwise known as phenyl-amido-propionic acid; R. Fittig and C. Wurster describe atropaic and isatropaic acid; and R. Fittig adds certain theoretical considerations on the formation of non-saturated hydrocarbons from the addition-products of the non-saturated acids.

Halogen Substitution-Products of Æthan.—W. Stadel.—The author describes the action of chlorine upon ethyl chloride, ethylen chloride, ethylen chloride, dichlorethyl chloride, and mono-chlorethylene chloride, adding the remark that he has studied penta-chlorethan, and is at present engaged with hexa-chlorethan, his results on both which bodies will shortly appear.

Chloro-brom- and Brom-Substitution-products of Ethylen.—Dr. J. Denzel.—The author describes the preparation and properties of α -chlor-brom-ethylen, α -chlor-dibrom-ethylen, α -dichlor-brom-ethylen, dichlor-dibrom-ethylen, and the brom-substitution-products of ethylen.

Nomenclature and Boiling-points of the Chloro-brom-substitution-products of Æthan and Ethylen.—Dr. J. Denzel.—The most important part of this paper is a table of boiling-points.

Simple Apparatus for Regulating and Varying the Atmospheric Pressure in Distillations, Determinations of Boiling-Points, &c.—W. Stadel and E. Hahn.

Reduction of Analytical Weighings to a Vacuum.—G. F. Becker.—These two papers require the accompanying plates. Becker lays down the law that the correction for the influence of the atmosphere decreases in proportion to the increase of the square of the specific gravity.

Experimental Researches on Hydrogen Peroxide.—E. Schöne.—The author examines the behaviour of hydrogen peroxide with potassium iodide, a point of great importance for ozonoscopic observations. The author concludes from numerous careful experiments that perfectly pure hydrogen peroxide, whether in the state of vapour or in aqueous solution, and whether in a concentrated or diluted condition, liberates iodine from potassic iodide.

Hexylens Formed from Tertiary Hexylic Alcohols and on their Polymerisation.—L. Jawein.—Not susceptible of useful abstraction.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 61, January, 1879.

Experiments Recently Undertaken to Dephosphorise Cast-iron.—M. Gruner.—The author has ascertained that dephosphorisation may be effected in the Bessemer apparatus or the Siemens furnace if the siliceous lining in both is dispensed with and refractory basic materials are used instead.

Metallurgy of Nickel.—As a note to Mr. A. H. Allen's paper translated from the *Journal of the Society of Arts* is appended an illustrated memoir by Badoureaux on the treatment of European ores of nickel and cobalt.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 6, 1879.

The Berlin Aniline Colour Company have issued a lengthy reply to the note by MM. Bindschedler and Bush on malachite green, inserted in the *CHEMICAL NEWS*, vol. xxxix., p. 61. They deny that a green dye can be industrially prepared from nitrobenzol and dimethyl-anilin by an oxidation process.

No. 7, February 13, 1879.

This issue contains a long account of the chemical manure works of M. Coquerel and Co., near Paris. This establishment, though only opened in 1874, sold, in 1877, six million kilos. of different manures.

MISCELLANEOUS.

The Royal Agricultural College, Cirencester.—The rumoured resignation of Prof. Church is incorrect. The Committee of Management of the Agricultural College are about to determine whether, after his marriage, he can retain his post without residing in the College precincts. Should this chair become vacant intending candidates may find the following extracts from the Collège Bye-Laws of some use:—

"XXI. The Principal . . . shall exercise supreme control over all Departments, Moral, Instructional, and Domestic, and over the Library, Museum, Gardens, Laboratory, and Veterinary Hospital.

"XXII. He . . . shall regulate the frequency and duration of the several Lectures and periodical Examinations, and all other matters of detail.

"XXIV. The Professors shall be appointed by the Principal, and be removable by him.

"XXVI. No private tuition by Professors is allowed, without the previous consent in writing of the Principal, to be separately given in every case.

"XXVII. The Professors shall be required to take a share in the maintenance of good order and discipline amongst the Students, subject to such regulations as the Principal shall appoint.

"XXVIII. All communications from the Professors and other officers of the College to the Council or Committee of Management shall be made only through the Principal."

The last Professor appointed has had to promise that he will engage in no literary work without leave from the Principal.

NOTES AND QUERIES.

Chromic Oxide.—Will any of your readers inform me how or where chromic oxide, as recommended by M. Audouin (*Les Mondes*, January 23, 1879) for fire-bricks, crucibles, &c., can be obtained at a cost that would allow of its being used for such a purpose.—CRUCIBLES.

MEETINGS FOR THE WEEK.

MONDAY, 10th.—Medical, 8.30.

Society of Arts, 8. "Dwelling Houses Their Sanitary Construction and Arrangements," by Dr. W. H. Corfield, M.A. (Cantor Lectures.)
London Institution, 5.
Royal Geographical, 1, 8.30.

TUESDAY, 11th.—Civil Engineers, 8.

Royal Institution, 3. "Animal Development," Prof. Schäfer.
Anthropological, 8.
Photographic, 8.

WEDNESDAY, 12th.—Society of Arts, 8. "The Compensation of

Watches, Clocks, and Chronometers," by E. Rigg, M.A.
Microscopical, 8.
Geological, 8.

THURSDAY, 13th.—Royal, 8.30.

Royal Institution, 3. "Sound," Prof. Tyndall.
Royal Society Club, 6.30.
Society of Arts, 8. "The Injurious Effects of the Air of Large Towns on Animal and Vegetable Life, and on Methods Proposed for Securing Salubrious Air," by W. E. Thomson, F.R.S.E.

FRIDAY, 14th.—Royal Institution, 9. "History of Games," by E. B. Tylor.

Quekett, 8.

SATURDAY, 15th.—Royal Institution, 3. "Colbert and Richelieu," by Mr. Walter H. Pollock.

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THE CHEMICAL NEWS.

VOL. XXIX. No. 1007

A NEW CHEMICAL INDUSTRY,
ESTABLISHED BY M. CAMILLE VINCENT.*

By Prof. ROSCOE, LL.D., F.R.S.

"AFTER I had made the discovery of the *marine acid air*, which the vapour of spirit of salt may properly enough be called, it occurred to me that, by a process similar to that by which this *acid air* is expelled from the spirit of salt, an *alkaline air* might be expelled from substances containing the volatile alkali. Accordingly I procured some volatile spirit of sal-ammoniac, and having put it into a thin phial and heated it with a flame of a candle, I presently found that a great quantity of vapour was discharged from it, and being received into a basin of quicksilver it continued in the form of a transparent and permanent air, not at all condensed by cold." These words, written by Joseph Priestley rather more than one hundred years ago, describe the experiment by which ammonia was first obtained in the gaseous state.

Unacquainted with the composition of this alkaline air, Priestley showed that it increased in volume when electric sparks are passed through it, or when the alkaline air (ammonia) is heated the residue consists of inflammable air (hydrogen).

Berthollet, in 1785, proved that this increase in bulk is due to the decomposition of ammonia into nitrogen and hydrogen, whilst Henry and Davy ascertained that two volumes of ammonia are resolved into one volume of nitrogen and three volumes of hydrogen.

The early history of sal-ammoniac and of ammonia is very obscure. The salt appears to have been brought into Europe from Asia in the seventh century, probably from volcanic sources. An artificial mode of producing the ammoniacal salts from decomposing animal matter was soon discovered, and the early alchemists were well acquainted with the carbonate under the name of *Spiritus salis urinae*. In later times sal-ammoniac was obtained from Egypt, where it was prepared by collecting the sublimate obtained by burning camels' dung.

Although we are constantly surrounded by an atmosphere of nitrogen, chemists have not yet succeeded in inducing this inert substance to combine readily, so that we are still dependent for our supply of combined nitrogen, whether as nitric acid or ammonia, upon the decomposition of the nitrogenous constituents of the bodies of plants and animals. This may be effected either by natural decay, giving rise to the ammonia which is always contained in the atmosphere, or by the dry distillation of the same bodies, that is, by heating them strongly out of contact with air; and it is from this source that the world derives the whole of its commercial ammonia and sal-ammoniac.

Coal, the remains of an ancient vegetable world, contains about 2 per cent. of nitrogen, the greater part of which is obtained in the form of ammonia when the coal undergoes the process of dry distillation. In round numbers two million tons of coal are annually distilled for the manufacture of coal gas in this country, and the ammoniacal water of the gasworks contains the salt of ammonium in solution.

According to the most reliable data 100 tons of coal were distilled so as to yield 10,000 cubic feet of gas of specific gravity 0.6, giving the following products, in tons:—

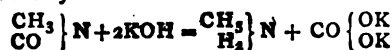
Gas.	Tar.	Ammonia Water.	Coke.
22.25	8.5	9.5	59.75 average.

* A Discourse given at the Royal Institution of Great Britain, Friday, February 21, 1879.

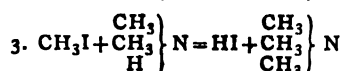
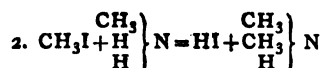
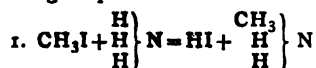
This ammonia water contains about 1.5 per cent. of ammonia; hence the total quantity of the volatile alkali obtainable from the gasworks in England amounts to some 9000 tons per annum.

A singular difference is observed between the dry distillation of altered woody fibre as we have it in coal, and woody fibre itself. In the products of the first operation we chiefly find in the tar the aromatic hydrocarbons, such as benzene, whilst in the second we find acetic acid and methyl alcohol are predominant.

The year 1848 is a memorable one in the annals of revolutionary chemistry, for in that year Wurtz proved that ammonia is in reality only one member of a very large family. By acting with caustic potash on the nitrates of the alcohol radicals he obtained the first series of the large class of compound ammonias the primar monamines, Of these methylamine is the first on our list:—



The years that followed, 1849 to 1851, were prolific in ammoniacal discoveries. Hofmann pointed out that not only one atom of hydrogen in ammonia can be replaced by its equivalent of organic radical, but that two or all the three atoms of the hydrogen in ammonia can be likewise replaced, giving rise to the secondary and tertiary amines, by the following simple reactions:—



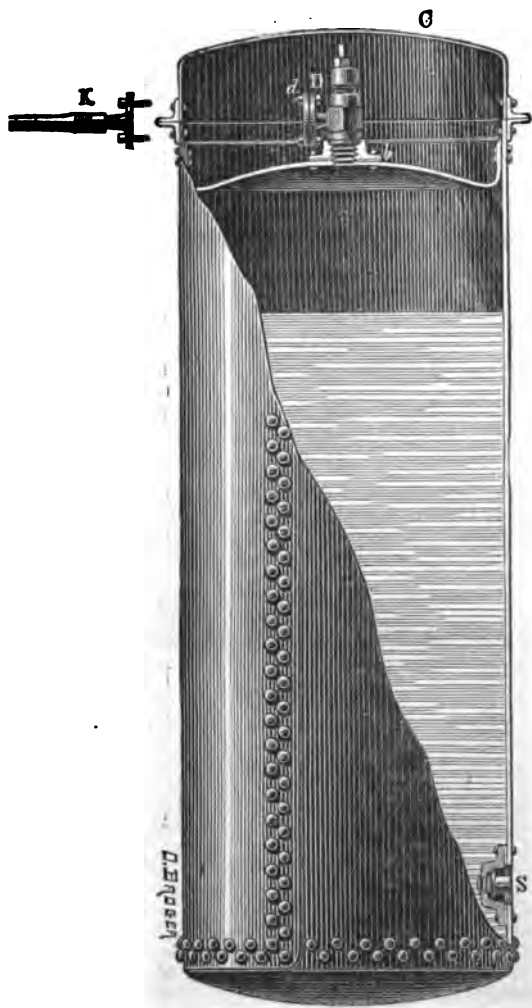
To these bodies the names of methylamin, di-methylamin, and tri-methylamin were given. They resemble ammonia in being volatile alkaline liquids or gases, which combine with acids to form crystalline and well-defined salts.

Hitherto these compound ammonias have been chemical curiosities; they have, however, recently become, as has so often been the case in other instances, of great commercial importance, and are now manufactured on a large scale.

We are all well aware that the French beet-root sugar industry is one of great magnitude, and that it has been largely extended in late years. In this industry, as in the manufacture of cane sugar, large quantities of molasses or treacle remain behind after the whole of the crystallisable sugar has been withdrawn. These molasses are invariably employed to yield alcohol by fermentation. The juice of the beet, as well as that of cane-sugar, contains, in addition to the sugar, a large quantity of extractive and nitrogenous matters, together with considerable quantities of alkaline salts. In some sugar-producing districts the waste-liquors or spent-wash from the stills—called *vinasses* in French—are wastefully and ignorantly thrown away, instead of being returned to the land as a fertiliser, and thus the soil becomes impoverished. In France it has long been the custom of the distiller to evaporate these liquors (*vinasses*) to dryness, and to calcine the mass in a reverberatory furnace, thus destroying the whole of the organic matter, but recovering the alkaline salts of the beet-root. In this way 2000 tons of carbonate of potash are annually produced in the French distilleries. For more than thirty years the idea has been entertained of collecting the ammonia-water, tar, and oils which are given off when this organic matter is calcined, but the practical realisation of the project has only quite recently been accomplished, and a most unexpected new field of chemical industry thus opened out, through the persevering and sagacious labours of M. Camille Vincent, of Paris.

The following is an outline of the process as carried out at the large distillery of Messrs. Tilloy, Delaune, and Co., at Courrières. The spent-wash having been evaporated until it has attained a specific gravity of 1.31, is allowed to run into cast-iron retorts, in which it is submitted to dry distillation. This process lasts four hours; the volatile products pass over, whilst a residue of porous charcoal and alkaline salts remains behind in the retort. The gaseous products given off during the distillation are passed through coolers, in order to condense all the portions which are liquid or solid at the ordinary temperature, and the combustible gases pass on uncondensed and serve as fuel for heating the retorts.

FIG. 1.



The liquid portion of the distillate is a very complex mixture of chemical compounds, resembling in this respect the corresponding product in the manufacture of coal-gas. Like this latter, the liquid distillate from the spent-wash may be divided into—

1. The ammonia water.
2. The tar.

The ammonia water of the vinasse resembles that of the coal-gas manufacture in so far as it contains carbonate, sulphurate, and hydrocyanide of ammonia; but it differs from this (and approximates to the products of the dry

distillation of wood) by containing in addition methyl alcohol, methyl sulphide, methyl cyanide, many of the members of the fatty acid series, and, most remarkable of all, *large quantities of the salts of trimethylamin.*

The tar, on re-distillation, yields more ammonia water, a large number of oils, the alkaloids of the pyriden series, solid hydrocarbons, carbolic acid, and, lastly, a pitch of fine quality.

The crude alkaline aqueous distillate is first neutralised by sulphuric acid, and the saline solution evaporated, when crystals of sulphate of ammonia are deposited; and these, after separating and draining off, leave a mother-liquor, which contains the more soluble sulphate of trimethylamin. During the process of concentration, vapours of methyl alcohol, methyl cyanide, and other nitriles are given off, these being condensed, and the cyanide used for the preparation of ammonia and acetic acid by decomposing it with an alkali.

Trimethylamin itself is at present of no commercial value, though perhaps the time is not far distant when an important use for this substance will be found. The question arises as to how this material can be made to yield substances capable of ready employment in the arts. This problem has been solved by M. Vincent in a most ingenious way. He finds that the hydrochlorate of trimethylamin, when heated to a temperature of 260°, decomposes into (1) ammonia, (2) free trimethylamin, and (3) chloride of methyl.



By bubbling the vapours through hydrochloric acid the alkaline gases are retained, and the gaseous chloride of methyl passes on to be purified by washing with dilute caustic soda and drying with strong sulphuric acid. This is then collected in a gas-holder, whence it is pumped into strong receivers and condensed.

The construction of these receivers is shown in Fig. 1. They consist of strong wrought-iron cylinders, tested to resist a pressure of 20 kilos. per square centimetre, and containing 50, 110, 220 kilos. chloride of methyl. The liquid is drawn from these receivers by opening the screw tap D, which is covered by a cap C, to prevent injury during transit.

Both ammonia and chloride of methyl are, however, substances possessing a considerable commercial value. The latter compound has up to this time, indeed, not been obtained in large quantities, but it can be employed for two distinct purposes: (1) it serves as a means of producing artificial cold; (2) it is most valuable for preparing methylated dyes, which are at present costly, inasmuch as they have hitherto been obtained by the use of methyl iodide, an expensive substance.

Methyl chloride was discovered in 1804 by MM. Dumas and Péligot, who obtained it by heating a mixture of common salt, methyl alcohol, and sulphuric acid. It is a gas at the ordinary temperature, possesses an ethereal smell and a sweet taste, and its specific gravity is 1.738. It is somewhat soluble in water (about 3 vols.), but much more in acetic acid (40 vols.), and in alcohol (35 vols.). It burns with a luminous flame, tinged at the edges with green, yielding carbonic and hydrochloric acids. Under pressure methyl chloride can be readily condensed to a colourless, very mobile liquid, boiling at -23°C. under a pressure of 760 m.m. As the tension of the vapour is not high, and as it does not increase very rapidly with the temperature, the liquefaction can be readily effected, and the collection and transport of the liquefied chloride can be carried on without danger.

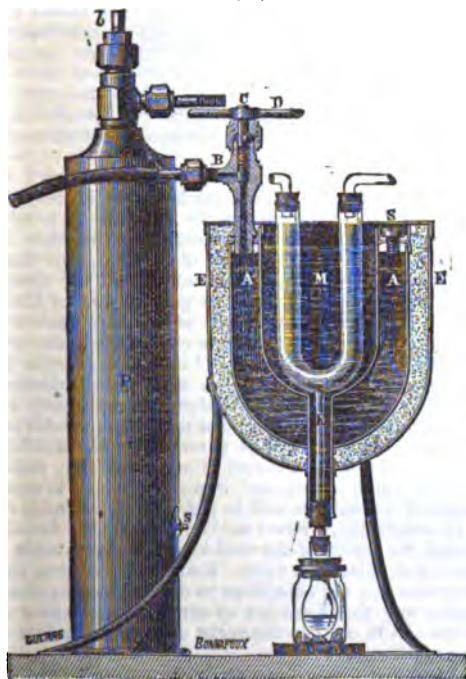
The following table shows the tension of chloride of methyl at varying temperatures:—

At 0° the tension of CH_3Cl is 2.48 atmospheres.				
" 15°	"	"	4.11	"
" 20°	"	"	4.81	"
" 25°	"	"	5.62	"
" 30°	"	"	6.50	"
" 35°	"	"	7.50	"

From these numbers we must of course subtract 1 to obtain the pressure which the vapour exerts on the containing vessel.

As a means of producing low temperatures chloride of methyl will prove of great service both in the laboratory and on a larger industrial scale. When the liquid is allowed to escape from the receiver into an open vessel, it begins to boil, and in a few moments the temperature of the liquid is lowered by the ebullition below -23° , the boiling-point of the chloride. The liquid then remains for a length of time in a quiescent state, and may be used as a freezing agent. By increasing the rapidity of the evaporation by means of a current of air blown through the liquid, or better by placing the liquid in connection with a good air-pump, the temperature of the liquid can in a few moments be reduced to -55° , and large masses of mercury easily solidified. The construction of a small freezing machine employed by M. Camille Vincent is shown in Fig. 2. It consists of a double-cased copper vessel, between the two casings of which the

FIG. 2.



methyl chloride (A) is introduced. The central space (M) is filled with some liquid such as alcohol, incapable of solidification. The chloride of methyl is allowed to enter from the cylindrical reservoir by the screw tap (B) and the screw (S) left open to permit of the escape of the gas. As soon as the whole mass of liquid has been reduced to a temperature of -23° , ebullition ceases, the screw (S) may be replaced, and if a temperature lower than -23° be required the tube (A) placed in connection with a good air-pump. By this simple means a litre of alcohol can be kept for several hours at temperatures either of -23° or -55° , and thus a large number of experiments can be performed, for which hitherto the expensive liquid nitrous oxide or solid carbonic acid was required.

M. Vincent has recently constructed a much larger and more perfect and continuous form of freezing machine, in which by means of an air-pump and a forcing-pump the chloride of methyl is evaporated in the freezing machine and again condensed in the cylinders. This enlarged form of apparatus will probably compete favourably with the ether, and sulphurous acid, freezing machines now in

use, as they can be simply constructed, and as the vapour and liquid do not attack metal and are non-poisonous, and as the frigorific effects which it is capable of producing are most energetic.

The second and perhaps more important application of methyl chloride is to the manufacture of methylated colours.

It is well known that rosanilin or aniline-red, $C_{20}H_{19}N_3$, yields compounds possessing a fine blue, violet, or green colour, when a portion of the hydrogen has been replaced by the radicals methyl or ethyl, and the larger the proportion of hydrogen replaced the deeper is the shade of violet which is produced. Thus we have triethyl rosanilin or Hofmann's* violet, $C_{20}H_{16}(C_2H_5)_3N_3$.

By replacing one or two atoms of the hydrogen of aniline by methyl and by oxidising the methyl anilines thus obtained, Charles Lauth obtained fine violet colours, whilst about the same time Hofmann observed the production of a bright green colouring matter, now known as iodine green, formed during the manufacture of the violet, and produced from the latter colour by the action of methyl iodide.

In order to prepare aniline-green from the pure chloride of methyl, a solution of methyl aniline-violet in methyl alcohol is placed in an iron digester and the liquid rendered alkaline by caustic soda. Having closed the digester, a given quantity of liquid chloride of methyl is introduced by opening a tap, and the digester thus charged is placed in a water-bath and heated by a jet of steam, until the temperature reaches 95° , and the indicated pressure amounts to from 4 to 5 atmospheres. As soon as the reaction is complete the hot water is replaced by cold, and the internal pressure reduced by opening the screw tap of the digester. The product of this reaction heated and filtered, yields the soluble and colourless base, whose salts are green. To the acidulated solution a zinc salt is added to form a double salt, and the green compound is then precipitated by the addition of common salt. By adding ammonia to a solution of the green salt, a colourless liquid is obtained, in which cloth mordanted with tannic acid and tartar emetic becomes dyed of a splendid green.

If rosanilin be substituted for methyl aniline in the preceding reaction Hofmann's violet is obtained. The application of methyl chloride to the preparation of violets and greens is, however, it must be remembered, not due to M. Vincent; it has been practised for some years by aniline-colour makers. M. Vincent's merit is in establishing a cheap method by which perfectly pure chloride of methyl can be obtained, and thus rendering the processes of the manufacture of colours much more certain than they have been hitherto.

The production of methyl violet from dimethyl aniline may be easily shown by heating this body with a small quantity of chloral hydrate, and then introducing some copper turnings into the hot liquid. On pouring the mixture into alcohol, the violet colour is well seen.

In reviewing this new chemical industry of the beet-root vinasses, one cannot help being struck by the knowledge and ability which have been so successfully expended by M. Camille Vincent on the working out of the processes.

Here, again, we have another instance of the utilisation of waste chemical products and of the preparation on a large scale of compounds hitherto known only as chemical rarities.

All those interested in scientific research must congratulate M. Camille Vincent on this most successful issue of his labours.

The Spottiswoode Testimonial.—The Committee appointed to receive subscriptions to present a bust of Mr. William Spottiswoode, Pres. R.S., to the Royal Institution as a testimonial of his valuable services as its Treasurer and Secretary successively, have engaged Mr. Richard Belt as the sculptor.

* Hofmann, *Proc. Roy. Soc.*, xiii., 13, 1863.

NOTE ON
VICTOR AND CARL MEYER'S NEW METHOD OF
DETERMINING VAPOUR-DENSITIES.

By GREVILLE WILLIAMS, F.R.S.

A GREAT number of processes for determining vapour-densities have been published during the last few years, but by far the greater number of them have only been used by their inventors, and none of them have entirely superseded the processes of Gay-Lussac and Dumas for determinations at ordinary pressures.

The last process of Victor and Carl Meyer* has, however, had a different fate, and from the moment it was published excited the earnest attention of chemists and physicists. In simplicity and accuracy it leaves nothing to be desired, and it possesses the great merit of being available at all temperatures up to the softening point of glass; and, doubtless, if the apparatus were constructed in metal its range of usefulness would be still further increased. In fact, MM. Victor and Carl Meyer may be congratulated upon having devised a method of determining vapour-densities which will probably supersede all those in use at the present time for ordinary pressures.

Amongst its other merits it can be constructed by any one with the greatest ease with the materials which are to be found in every laboratory.

Having to determine the vapour-densities of two fractions of β -lutidine, supposed to be very nearly pure, I improvised an apparatus for the purpose. In all essential features it resembled that of Victor and Carl Meyer. The glass flask *c*, in the illustration, p. 66, was, however, replaced by an iron tube closed at the lower end, which happened to be in the laboratory. It was charged with aniline. The long neck of the vapour flask, *b*, was cut off two inches below the lateral tube, *a*, and, the two ends being brought close together, were joined by an india-rubber tube; the object being to give the apparatus some flexibility, and lessen the chances of fracture of the tube *a*, which had been drawn out somewhat too finely. These details are merely mentioned to show that, as long as the essential points of the apparatus are preserved, some latitude may be permitted in putting it together. In my first apparatus the funnel *d* was made as directed, but subsequent experience has shown that it is unnecessary, as the small tube is easily closed by an india-rubber stopper without the necessity for enlarging the aperture.

A preliminary experiment was made with water with the following result:—

$$\begin{aligned} S &= 0.0207 \text{ grm.} \\ t &= 13.1^\circ \\ B &= 758.8 \text{ m.m. (corrected.)} \\ w &= 11.1 \\ V &= 26.5 \text{ c.c.} \\ D &= 0.643 \end{aligned}$$

Experiment.	Theory.	Error.
0.643	0.622	+0.021

Two vapour-densities were then taken with β -lutidine—

Boiling-point, 164° .	Boiling-point, 164° to 165° .
$S = 0.0894$	0.0693
$t = 12^\circ$	12°
$B = 761.5 \text{ m.m. (corr'd.)}$	761.5
$w = 10.2$	10.2
$V = 20.0 \text{ c.c.}$	15.5
$D = 3.65$	3.65

Experiment.	Theory.	Error.
3.65 3.65	3.699	-0.049

I attribute part even of this small error to the β -lutidine containing traces of the next lower homologue.

* *Dent. Chem. Ges. Ber.*, xi., 2253; *CHEMICAL NEWS*, vol. xxxix. page 66.

ON THE ACTION OF NUCLEI, &c.

By C. TOMLINSON, F.R.S.

My paper to which Mr. Grenfell refers (*CHEMICAL NEWS*, vol. xxxix., p. 16) was not read in September, 1877, and its object was not "to upset his theories and establish my own." Is it not possible for a scientific man to be anxious only for the truth? That I was so will appear from the fact that Mr. Grenfell showed me some of his experiments, and although I did not agree with his conclusions, while he decidedly opposed mine, I asked him to draw up an account of them for the Royal Society, and I promised to present his paper, which I accordingly did, and it was read and printed in the *Proceedings*.

My object in repeating Mr. Grenfell's experiments was to see whether any fresh hint could be gathered from them. I never once mentioned my own theory; but I gained this useful piece of information, viz., "that supersaturated saline solutions behave differently among themselves and to different bodies under varying atmospheric conditions;" and that "an oil, &c., introduced into the closed flask, as in M. Viollette's and Mr. Liversidge's experiments, may be inactive, while under the other conditions the same oil may be active." I give abundant proof that this is so in a subsequent paper, which Mr. Grenfell does not appear to have seen; while in a third paper the powerful influence of the sides of the vessel in maintaining the state of supersaturation is dwelt upon.

As space is valuable in the *CHEMICAL NEWS* I will not follow Mr. Grenfell in his method of minute criticism, which throws no light upon the question as to what are the precise conditions under which these solutions become solid. Mr. Grenfell says that it is absorption that does the whole work. That is a fair subject for discussion; but I must protest against the general tone of Mr. Grenfell's note, especially that part of it which makes me so silly as not to know the simple conditions under which he conducted his experiments. In such a censure he, unconsciously perhaps, includes such good observers as Löwel, Viollette, and Liversidge, who expressly say that porous bodies, bodies greedy of water and capable of being hydrated, are incapable of determining the solidification of these solutions. I stated in one of my early papers that I do not agree to this, and in a paper which is now being prepared the reasons will be given more distinctly.

In conclusion, I never said or implied that Mr. Grenfell denied the result of the well-known lecture table experiment with Glauber's salts. I may also add that the early observers saw the objections to cotton-wool for closing the flasks, and that I am not so careless as to allow crystals of the salt to exist in the necks of my flasks before making an experiment.

Highgate, N., March 8, 1879.

LARGE CRYSTAL GROWING.

By CHARLES W. QUIN.

WHEN I was a student I was much given to large crystal growing. My great difficulty was to guard against sudden rises in temperature, which generally had the effect of causing the growing crystal to be partially and unevenly re-dissolved. To receive a continually even temperature, after many experiments I hit upon the plan of plunging the beaker containing the growing crystal into the house-cistern, the temperature of the water in which, as I found by trial, never differed by more than 0.5°F. either way day or night. To prevent the solution from becoming exhausted too rapidly I used to immerse it to the depth of half an inch or so a crystal drainer containing a filter-paper full of the salt to be crystallised, so that a constant stream of strong solution was continually descending on the growing crystal.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, March 6, 1879.

Dr. J. H. GLADSTONE, F.R.S., President, in the Chair.

AFTER the announcement of visitors, confirmation of minutes, &c., the following certificates were read for the first time:—F. A. B. Jewson, T. H. Walker, W. E. Blythe. The list of Officers proposed by the Council was then read from the Chair.

The PRESIDENT then called on Mr. G. ATTWOOD to read a paper "*On the Quantitative Blowpipe Assay of Mercury.*" The author divides compounds to be assayed into three classes. Class A, containing metallic mercury, cinnabar, tiemannite, suboxide, protoxide, and mixed sulphides. Class B, calomel, corrosive sublimate, and iodide of mercury. Class C, amalgams of gold, silver, copper, lead, zinc, tin, &c. Class A.—10 to 20 grains of the ore, finely powdered and passed through a sieve, 2000 holes to the linear inch, are mixed with 5 to 10 times their weight of powdered litharge and distilled over a spirit-lamp in a small glass retort, $1\frac{1}{4}$ inches long and $\frac{1}{2}$ inch in diameter. To this retort is fitted by means of a cork a glass tube, slightly curved, $2\frac{1}{2}$ inches long, and $\frac{1}{4}$ ths of an inch in diameter. The end of this tube dips under water contained in a small porcelain crucible. The operation lasts only a few minutes. The mercury is carefully collected from the glass tube and crucible. The retort is broken up and its contents carefully powdered and examined by a lens for mercury. The globules are then united by gently warming under water, and the dry mercury weighed. Class B.—A quantity of the finely powdered ore, equal to 10 grs., is mixed with three times its volume of oxalate of potash and one volume of cyanide of potassium. The apparatus closely resembles that used in class A, but the retort has a small bulb. Class C.—These amalgams are sometimes powdered with difficulty, and it is often advantageous to add a known weight of pure mercury, so as to render them semifluid before distilling. 10 to 30 grs. of the amalgam are usually taken for an assay. A turned steel retort is used for distillation, which is effected in a small charcoal furnace heated by a blowpipe flame; the head of the retort is accurately ground to fit over the body. The retort including the cup and cap is 1 inch high; the neck of the cap is 2 inches long. The paper contains full-size illustrations of the different retorts, &c., which are made by Casella. The author has had much experience, and states that most accurate results can be obtained with the above apparatus.

The next paper was read by Mr. J. W. THOMAS. "*On some Points in the Analysis of Combustible Gases and in the Construction of Apparatus.*" In 1874 the author noticed that when a small quantity of marsh gas was mixed with about three times its volume of oxygen and rarefied until the pressure was 160 m.m. no explosion took place; similarly when mixed with twice its volume of oxygen under a pressure of 130 m.m. the spark did not ignite the mixture. It soon became evident that the cooling effect of the walls of the eudiometer was the chief agent in modifying the force of the explosion. It was also noticed that the expansion of a gas to twice its volume lessened the force of the explosion to a greater extent than the addition of an equal volume of inert gas at the initial pressure. The author accordingly made estimations with marsh gas and hydrogen, using nearly the theoretical quantities of oxygen and a pressure of 160 to 170 m.m., and found that perfectly accurate results were obtained, whilst the safety of the eudiometer tube was not in the least endangered. The author then proceeds to point out the errors which creep in when the long (800 to 900 m.m.) eudiometer tube of Frankland and Ward's or

McLeod's apparatus is used, and proves that the sensitiveness of the apparatus depends on the length of the pressure tube above the top of the eudiometer tube. With the old method of introducing a large excess of oxygen in order to moderate the violence of the explosion, this long eudiometer tube was necessary to contain the large quantity of gas introduced. On the other hand, to lengthen the pressure tube would render it unwieldy from its great length, so that with the old method of introducing a large excess of oxygen the relative lengths of the pressure and eudiometer tubes must remain unaltered; but by employing the reduced tension method proposed by the author, and in which only the theoretical quantity of oxygen required has to be introduced, the necessity for such a long eudiometer tube is obviated. The author, therefore, has shortened the eudiometer tube to about 500 m.m., retaining the original length of the pressure tube, and finds that the apparatus has gained in delicacy and is still sufficiently large for all substances. The next modification introduced by the author is the substitution of a steel block with a three way steel tap for the glass taps connecting the barometer tube, eudiometer tube, and pressure bottle. The tubes are fixed and held tight in the steel block by india-rubber rings, screwed down by means of steel collars. Flexibility and absolute tightness are thus secured. As regards the steel tap the author has had no difficulty in keeping it perfectly tight; the three ways are gouged out on the surface of the tap and are not bored. The use of the steel plates connecting the eudiometer and laboratory tubes has been abandoned, and a hollow glass tap substituted, which is so bored that the tubes can be washed out. The mercury trough is made movable by an ingenious mechanical arrangement. The supply of water to keep the apparatus at a constant temperature is brought in at the top of the barometer tube; the exit is at the bottom, and a syphon arrangement is added to ensure a thorough mixing of the water round the top of the eudiometer tube. The little windlass has been modified so that it can be worked with one hand. In conclusion the author gives details as to the management of a gas analysis with the modified method and apparatus; a drawing of the latter accompanies the paper.

Prof. FRANKLAND complimented the author on the ingenuity displayed and the success achieved in his paper. At the first mention of the return to a use of the steel stopcock he must confess that he had almost shuddered. Twenty-five years ago when glass stopcocks were a luxury these steel taps constituted a never-ending source of annoyance. After a week's work a leak almost always occurred, which necessitated a prolonged and careful grinding. This might be due to the fact that the tap was made of cast-iron and the plug was bored. The method of exploding gases with almost theoretical quantities of oxygen was a decided step in advance. The shortening of the eudiometer tube and the increased sensitiveness thereby attained seemed to him also most important improvements. He would like to ask Mr. Thomas how the bursting of the flexible tube connected with the pressure bottle was obviated. All plans that he had tried had ultimately failed (winding tape, tubing with canvas in it, &c.).

Dr. WRIGHT suggested that the tubing should be coiled round with bell-wire.

Prof. McLEOD said that a plan which answered very well was to bind the tube with tape, and cover that with sheet india-rubber to prevent contact of the tape with potash, &c. He had been much struck with the success with which Mr. Thomas had used the theoretical quantity of oxygen without fracturing the eudiometer. In the old apparatus the difficulty was not so much to keep the steel tap tight as to make a tight cement joint between the block and the glass tubes.

Mr. HART suggested that the india-rubber tubing should be wound with tape soaked in glue containing bichromate of potash. On subsequent exposure to light a compound was formed unattacked by acids and alkalis.

Dr. ARMSTRONG had used a gas apparatus made after Mr. Thomas's model with great success. His stopcock was bored. The leakage of the old stopcocks might be due to corrosion by resin cerate. Bullocks' fat was most suitable for lubrication (Prof. Frankland stated that melted india-rubber was always used, not resin cerate, for the taps). He suggested that the mechanism employed by Mr. Thomas for shifting the mercury trough caused much vibration: he had used a small hydraulic lift. If the india-rubber tube were covered with double tape sewed longitudinally no difficulty was experienced.

Mr. THOMAS, in reply, pointed out that his steel tap was of a much larger size than the old one. In answer to Dr. Armstrong, no vibration was caused by the turning aside of the mercury trough.

The next paper was read by Mr. J. M. THOMSON, "*On the Action of Isomorphous Salts in Exciting the Crystallisation of Supersaturated Solutions of each other, and some Experiments on Supersaturated Solutions of Mixed Salts.*" In this paper the author communicates the results of more than 400 observations. The following are given as the general results to be deduced from the experiments. The results of Gernez (*Ann. Sci. de l'Ecole, norm. sup.*, 1861) have been confirmed and extended, showing that truly isomorphous bodies act as exciting nuclei in inducing the crystallisation of supersaturated solutions; that mere form alone does not render the body active to a supersaturated solution of a salt isomorphous with the nucleus; but that it is necessary for the nucleus to possess an identical chemical structure. Thus, cubes of iron pyrites and octahedral crystals of magnetite are inactive to alum solutions. In the case of mixtures of solutions two series of results were experienced in the same solution:—A. When the mixture consists of two salts not isomorphous—(1) Sudden crystallisation may commence, spreading, however, gradually through the solution, on the addition of a nucleus, causing a deposition of the body belonging to the nucleus only. (2) That when sudden crystallisation takes place, causing the deposition of both salts, there is a preponderance of the salt of the same nature as the nucleus. (3) That the nucleus may remain growing slowly in the solution, becoming increased by a deposition of the salt of the same nature as the nucleus. B. When a mixture consists of two isomorphous salts—(1) Sudden crystallisation may occur, giving a deposition of both salts apparently in the proportions in which they exist in solution. (2) That when slow crystallisation takes place the nucleus increases by a deposition of the least soluble salt, showing that in mixed supersaturated solutions a gradation of phenomena may be experienced passing from those shown in the crystallisation of a true supersaturated solution to those shown in the crystallisation of an ordinary saturated solution. The experiments were chiefly made as follows:—The saturated solution to be used as a nucleus was placed either in a bulb-tube or a tube bent at its lower end into a double U (as used by Liversidge). The solution was then boiled, and the tube plugged with cotton-wool. The supersaturated solution was introduced into a small flask. The tube with nucleus solution is plugged tightly into the neck of the flask, and the solution boiled. When cold, the nucleus solution is caused to crystallise, and lowered so that at first the glass only touches the solution; this produced uniformly negative results. On bringing the crystals in contact by breaking the bulb or lowering the U-tube crystallisation commenced if the nucleus was active. The author is continuing his experiments, and promises a further communication on the subject. The substances already experimented with include—Magnesium sulphate, $7\text{H}_2\text{O}$, to which zinc, nickel, cobalt, iron sulphates with $7\text{H}_2\text{O}$ are all active; sodium sulphate with 10 or $5\text{H}_2\text{O}$ being inactive; sodium sulphate and seleniate, $10\text{H}_2\text{O}$, are active to sodium sulphate solution, $10\text{H}_2\text{O}$. Chrome and iron potassium alums with $12\text{H}_2\text{O}$ are active to the ordinary potash alum, &c.

The PRESIDENT remarked how various were the conclusions which had been arrived at concerning the crystalli-

sation of supersaturated solution by different experimenters. Mr. Thomson's researches had modified the difficulties which existed as to the theory that the cause was the introduction of a crystal of the same salt, by pointing out that a similar salt would also act as a nucleus, and the method might prove an accurate means of determining whether two bodies of similar, were really of identical, chemical composition or not.

Dr. WITT, in reply to a question put by Mr. Thomson, said that supersaturated solutions were of frequent occurrence in the salts of the aromatic substances, disulpho-derivatives of benzene, the acetates of the different rosanilins, &c. He asked whether the phenomena of supersaturation were included by Mr. Thomson in the same class as those of supersaturation.

Prof. HARTLEY suggested that the crystallisation of one salt out of a mixture, by introducing a nucleus active to it but inactive to the other salts, might be very useful for separation on the manufacturing scale.

Mr. GROSJEAN said that Crace-Calvert had thus crystallised carbolic acid from a mixture of that substance and cresolic acid.

Mr. THOMSON did not think any strict line could be drawn between supersaturation and supersaturation.

The SECRETARY then read a paper by WATSON SMITH, "*On the Isomeric Dinaphthyls.*" This communication is a continuation of the author's previous work. The immediate objects of the present investigation were—(1) To prepare a quantity of the α - α -dinaphthyl (melting-point 154°) by Lossen's method, in order to compare it with the body previously obtained by the author in yellowish plates (m.p. 147°). (2) To examine the crude product obtained by Lossen to see if it contains besides the α - α -dinaphthyl an appreciable quantity of the other two dinaphthyls. (3) To prepare sufficient quantities of the three isomers in a completely pure condition, and to determine their vapour-densities. (4) To discover further reactions in which dinaphthyl is formed, so as to find out, if possible, so much about the conditions of its formation as would lead to a method of preparation more easily carried out and giving a larger yield of product than the present method. (1) The author prepared the α - α -dinaphthyl, and found that it melted at 147° , but by further boiling with animal charcoal in petroleum spirit, colourless crystals were obtained, melting at 154° to 155° . (2) The β - β - and the α - β -dinaphthyls could not be detected. (3) The pure substances were obtained, and their vapour-densities determined by V. Meyer's new apparatus with great facility; a lead-bath and nitrogen atmosphere were used. β - β -dinaphthyl, m.p. 187° , v.d. 8.73; α - α , m.p. 154° , v.d. 8.67; α - β , m.p. 76° , v.d. 8.78; calculated v.d., 8.77. In the fourth part the author has investigated the action of carbon tetrachloride, chloroform, carbon disulphide, bromo-naphthalene on naphthalene. He has further investigated the reactions which take place when a mixture of bromo-naphthalene and naphthalene are passed over heated soda-lime, lime, ferric oxide, and silver.

The Society then adjourned to March 20, when the following papers will be read:—"On Perplumbic Ethide," by E. Frankland and A. Laurance; "On the Decomposition of Water by certain Metalloids," by C. F. Cross and A. Higgin; "On the Volumetric Determination of Chromium," by W. J. Sole; "The Production of the Higher Oxides of Iron, Chromium, Manganese, and Bismuth," by W. Foster.

PHYSICAL SOCIETY.

Ordinary Meeting, March 8, 1879.

Prof. W. G. ADAMS, President, in the Chair.

Dr. HURST and Mr. Jacob were elected Members.

Prof. AYRTON brought forward a new theory of terrestrial magnetism, originated by himself and Prof. Perry, of the Imperial Engineering College, Japan. It is well known

that metal cages act as screens against induction in the case of static electricity or electricity at rest, and hence Clerk Maxwell, at the British Association meeting for 1876, suggested that no earth connection was necessary for lightning-conductors, since a cage would be sufficient. But dynamic electricity is different from static in this respect, and Profs. Ayrton and Perry found that even a thick block of copper will not screen a coil of wire from the induction by a current flowing in a neighbouring one. Some experiments of Dr. Muirhead, not yet published, would seem to favour the view that a current is a series of intermittent changes of potential, and that the inductive effect was due to a difference in the epochs of the currents in the two coils. It was found by Helmholtz that a quantity of static electricity in mechanical motion performs work conversely. Mr. Crookes finds that the stream of molecules from a -pole *in vacuo* is electrified and may be deflected by a magnet. It is upon that fact that Profs. Ayrton and Perry have based their theory, which is easily explained by supposing the earth to be an isolated sphere with a static charge residing on its surface. Then, since each electrified particle at the surface will be moving relatively to a point in the interior, it follows that the interior must be magnetic. The theory is independent of the substance of the interior; but in order to simplify the working the authors treated the case of a solid iron ball, and curiously enough arrived at the result expressed by Biot's law for the distribution of magnetism on the surface of the earth:—

$$I = M \sqrt{1 + 3 \cos^2 \theta}.$$

And, similarly, they found that if the earth were electrified to the potential of 10^6 volts. relatively to interplanetary space, its magnetisation would be as it is. If the earth were alone in the universe, then, by this theory, it would have its own magnetic state by virtue of its electric charge and axial rotation. If other bodies in the universe, however, had their magnetic states too, these would influence the earth's, and hence we should have terrestrial tides and storms of magnetic force, such as are known to exist, as, for instance, when changes take place in the sun's atmosphere by approach of planets and other causes. Lastly, the iron in the interior of the earth may give it a certain amount of coercive force, but the theory does not rest on this.

Dr. J. HOPKINSON then read an account of some experiments with the quadrant electrometer, which showed that Clerk Maxwell's formula for the sensibility of the electrometer,—

$$(A - B) \left(C - \frac{A + B}{2} \right),$$

where A and B are the potentials of the two pairs of quadrants, and C the potential of the needle, only holds good when C (the charge of the jar or needle) is less than 200 Daniell elements. Above that a different law appears to hold. Dr. Hopkinson also remarked that any degree of low sensibility down to zero could be got from the electrometer by connecting a condenser to each pair of quadrants, and adjusting their capacities.

Mr. F. D. BROWN described his apparatus for maintaining constant temperatures and pressures. A constant temperature can be obtained if the pressure can be kept constant. The vessel in which the constant pressure is desired communicates with an air-pump by a pipe in which a movable tap or valve is placed. By opening or closing this tap the pressure is regulated. This is effected by an electric clutch arrangement. A mercury anemometer sends a positive or negative current from a battery through the clutch according as the pressure is too high or low, and this current actuates the clutch to close or open the valve. The clutch consists of an axle driven by a turbine to get power to work the valve, and the current, by means of electro-magnetism, connects the tap to the axle, which then opens or closes it as the case may be. In this way a pressure varying no more than one-fifth millimetre each way can be obtained.

MANCHESTER LITERARY AND PHILOSOPHICAL SOCIETY.

Ordinary Meeting, February 18, 1872.

J. P. JOULE, D.C.L., LL.D., F.R.S., &c., President, in the Chair.

JAMES BOTTOMLEY, B.A., D.Sc., and RICHARD S. DALE, B.A., were appointed Auditors of the Treasurer's accounts. "On a Chemical Investigation of Japanese Lacquer, or 'Urushi,'" by SADAMU ISHIMATSU. Communicated by Professor ROSCOE, LL.D., F.R.S.

During a few months last year I had the opportunity of examining roughly into the nature of "Urushi," in the Laboratory of Tokio University.

The specimen of lacquer which I had under my examination was obtained from Kuyemon Nakamuraya, in Tokio, a large lacquer merchant.

It is a milky juice of pale grey colour, and gives out a certain kind of poisonous volatile gas. Some persons are terribly attacked by this poison, producing a great swelling where the acid comes in contact. During my examination in the laboratory one of the apparatus keepers was terribly attacked by this gas, producing ugly swellings all over the face. He told me at the time it was exceedingly itchy. By using the solution of chloride of sodium, carbonate of soda, acetate of lead, &c., he was said to have recovered within a week. This poison acts only on certain persons. I had to work with it for many days, yet never had any attack of the kind nor felt any uneasiness from it.

It has a sweetish characteristic smell, and has an irritating taste. It burns with very luminous flame, evolving dense black smoke like oil of turpentine.

It is to a great extent soluble in benzol, ether, absolute alcohol, &c., leaving behind a blackish grey residue in which gum was found.

Lacquer on exposure to the atmosphere rapidly loses its weight, and at the same time blackens on its surface; although this loss is different in different specimens, yet on the average of those which I have examined it seems to vary from 25 to 37 per cent.

When the lacquer is exposed to the action of sunlight in hermetically sealed vessels in the atmosphere or in carbonic acid, blackening does not take place, but a large quantity of moisture collects on the sides of the vessel.

The loss of weight in the atmosphere is almost if not entirely due to the escape of water, with a minute quantity of carbonic acid which may be formed by the oxidation of some organic compound existing in the lacquer. The attempt has been made to estimate the relative amounts of carbonic acid and water; yet it was not successful at the time, being too difficult, and it must be left open to some future investigation.

It is by many supposed to be due to the combined action of light and air that the blackening of lacquer in the air takes place; but this seems to be erroneous from the following experiments:—First. I made a square box which had a well fitting sliding door, and the inside of which was made perfectly black, so that practically no light is permitted to enter. In it was placed a small quantity of lacquer at dark, and the door closely shut. On looking at it the next morning it was observed that the lacquer had turned perfectly black, proving that it is not the light that blackens the lacquer.

Second. The bottle in which I kept my lacquer for more than three months during my examination was exposed to the incident light of the laboratory. The surface of the lacquer was turned perfectly black, forming a wall as it were; while those portions which were in contact with the sides of the bottle, which receive as much light as if there were not any glass sides before it, were not at all blackened. This phenomenon is just complementary to the first one, proving that the blackening in the atmosphere is in all probability due to the oxygen of the air, but not the light alone, nor the combined action of light and air.

The lacquer when distilled with water gives a colourless distillate which is slightly acid to test-paper, and the attempt has been made to examine the acid, but not successfully on account of too minute quantity of the substance evolved. Distillation by itself and in a current of steam was tried also, but the results in both cases were the same as the first one. Lastly, distilled with a small quantity of dilute sulphuric acid, to aid the substance to distil over, into sugar of lead, scarcely any precipitate was obtained.

Lacquer mixes with any kind of fixed oil in all proportions; hence oil is often added as adulteration, but sometimes it is purposely added to increase its mobility.

The specimen of lacquer which I examined consisted of the following three substances:—

	I.	II.
Part soluble in absolute alcohol (resin)	58.24	58.23
Gum	6.34	6.30
Residue	2.24	2.30
Moisture and other volatile matter	33.175	33.170
	100.00	100.00

As I have already mentioned, the lacquer loses its weight very rapidly when exposed to the atmosphere. For the above determination I weighed out each time samples from well stoppered bottles, and determined the weight by difference. Then this was treated with absolute alcohol and the filtrate evaporated to small bulk and dried in an air-bath at 100° C. until the weight remained constant. This is put down as the part soluble in alcohol in the above analysis. The residue was then treated with hot water, and the filtrate evaporated and dried at 100° and weighed as gum. The residue, after the gum had been removed, was then washed, and dried on weighed filter at 100° C., and weighed as residue. The moisture and other volatile matter were of course determined by the difference.

The estimation of the amount of part soluble in alcohol, after the lacquer had been exposed to the sunlight in open vessel for some twenty or thirty days, shows that the soluble part increased up to 72.82 per cent. This number, when calculated for the substance to have lost 28 per cent moisture and other volatile matter during exposure, then equals 58.3 per cent, which is nearly equal and practically the same as the analysis previously given; hence there seems to have been no material change in the amount of matter soluble in alcohol. Now the perfectly dried lacquer, after being finely powdered, was dried at 100°, and analysis gave:—

Part soluble in absolute alcohol	18.07
Gum	3.63
Residue	78.30
	100.00

Altogether, from this analysis, the residue being increased, the lacquer seems to have undergone some change; but possibly this is owing to the fact that the alcohol as well as water seem to have had less complete access to the material.

Thus the "Urushi" consists of three principal constituents, (1) a resinous part soluble in alcohol, (2) gum, and (3) residue. Although there are, in addition to these, water and volatile matter, as they go away sooner or later before it is used they are not properly called the constituents.

(1) Part soluble in alcohol (resin) seems to be the principal portion, and has a smell like ordinary lacquer, but it never dries as the original does. It is brownish black, and slightly sticky to the touch. When treated with potash solution it forms a bluish black precipitate, but nothing is obtained on addition of sulphuric acid to the filtrate.

When boiled with hydrochloric acid the resin is transformed into a substance elastic while hot, something like the mass obtained when heated sulphur is dropped into

cold water. When boiled with nitric acid nitrous fumes are given out, and the mass gradually becomes yellow, and finally a beautiful orange coloured mass was obtained. This mass was washed with hot water several times, and then treated with absolute alcohol. The mass was to a great extent soluble in the alcohol, leaving behind a small quantity of a yellowish body (which I think to be part not sufficiently acted upon by the acid). This alcoholic extract forms a beautiful yellow precipitate with both nitrate of silver and acetate of lead. I took a quantity of alcoholic extract, precipitated it with acetate of lead, and the precipitate was thoroughly washed with absolute alcohol, and then decomposed by means of dilute sulphuric acid. (It might be better to decompose this salt with sulphuretted hydrogen, but we cannot do so on account of reducing action of this gas.) The mass was dissolved again in absolute alcohol, then separated from sulphate of lead.

Now then this separated alcoholic extract was again precipitated by sugar of lead, and, after filtering and washing, the precipitate was dried partially in an air-bath, and carried under the receiver of an air-pump and dried over sulphuric acid.

This lead salt exploded when heated. The amount of lead was estimated as oxide by igniting it with nitric acid, and the salt was subjected to organic combustion. Nitrogen was determined by Dumas's method.

The following numbers were obtained as the mean results:—

Carbon	26.93
Hydrogen	4.11
NO ₂	18.44
PbO	47.42
Oxygen	3.10
	100.00

Now I prepared the silver salt of the same, and obtained 18.5 per cent of silver as the result. It seems to give no help as to the formula of this body.

As such was the case, I took alcoholic extract of the original lacquer and precipitated it with acetate of lead, and after requisite purification and drying the precipitate was analysed. Lead determined as before.

The following is the mean of two experiments which I was only able to try:—

Carbon	50.450
Hydrogen	5.705
PbO	3.775
Oxygen	40.070
	100.000

(2) The gum is soluble in cold as well as in warm water. It has no smell, almost no taste; it has a yellowish or rather brownish colour, and is of a non-crystalline body. It is quite insoluble in alcohol.

On subjecting this substance to organic analysis I got the following percentage of hydrogen, carbon, and oxygen:—

	I.	II.
Carbon	41.20	41.45
Hydrogen	6.51	6.58
Oxygen	52.29	51.97
	100.00	100.00

These analyses yield a formula approximating to the composition of common gum.

(3) The residue is, I think, nothing more than the mixture of cellulose, bark, dust, &c.

In concluding my paper I must say that I am not at all satisfied with my present analyses, but I thought it might be of some interest to some of you from the point that, although the varnished articles from this juice are so celebrated, yet, as far as I am aware, this is the first analysis of the kind that has been heretofore attempted, and might be of some use to those who are interested upon this subject.

CORRESPONDENCE.

THE INSTITUTE OF CHEMISTRY.

To the Editor of the Chemical News.

SIR,—Judging from the erroneous statements contained in a letter signed "A Disgusted Promoter," in the CHEMICAL NEWS, vol. xxxix., p. 102, it would seem either that all the notices and papers sent to Members had miscarried in his case, or that the writer, although on the original Committee, as he intimates, is not actually a Fellow of the Institute.

So far from it being true "that the executive dared not publish a report of the meeting because nearly all the speakers upheld the system of giving certificates for advertising purposes," a report of the Conference on Trade Certificates has been printed for the use of Members. Again, the writer says—"In a few days' time a second conference will be held, at which the subject of the 'Adulteration of Food and Drugs' will be discussed." If he were actually a Fellow of the Institute he ought to know that that conference took place a fortnight ago, and that the notice sent out stated the subject to be "Adulteration of Articles of Food," and not "Adulteration of Food and Drugs." If, however, "A Disgusted Promoter" is really a Fellow of the Institute, and through some extraordinary and unfortunate coincidence all the notices, &c., have miscarried in his case, I should feel much obliged if he would communicate with me on the subject, and I will endeavour to ascertain the cause.

In conclusion, I may say that although the Council welcome open and fair criticism of their proceedings by independent Members, an anonymous letter recommending the Fellows not to pay their subscriptions can scarcely be considered as of that nature.—I am, &c.,

CHARLES E. GROVES, Secretary.

Somerset House Terrace, W.C.

CRYSTALLISATION OF PHOSPHORUS.

To the Editor of the Chemical News.

SIR,—Seeing Mr. Whewell's interesting contribution on farina in the CHEMICAL NEWS (vol. xxxix., p. 97) reminds me of a question I have often wished to ask. Having been now interested for some years in the crystallisation of phosphorus, anthracene, &c., *in vacuo*, I have eagerly looked for further details of Mr. Whewell's experiments (vol. xxviii., p. 205). Mr. Whewell there states that if the phosphorus is heated and spread over the sides of the tube "beautiful colourless crystals" will be obtained, "which will be slightly red." He also states that having made some of these peculiar crystals he sent them to a well-known mineralogist, who promised to examine them. Is Mr. Whewell in a position now to give us more information concerning them or even the well-known mineralogist's report.—I am, &c.,

GEORGE E. DAVIS.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 7, February 17, 1879.

Researches on Electric Fishes; Characters of the Discharge of the Gymnotus; Effects of the Discharge of the Torpedo Received in a Telephone.—E. J. Marey.—Physiologists have been struck with the analogies presented by a muscle and the apparatus of electric fishes. These two kinds of organs, both subject

to the will and provided with nerves for centrifugal action, have further a very analogous chemical composition, and even some points of structural resemblance. These views, put forward before physicists had developed the theory of the correlation of forces, were necessarily very vague. It might even be conceived that in the living organism, as well as in our physical instruments, analogous conditions might produce either mechanical work or electricity. Having found that muscular acts are complex, *i.e.*, that a muscle in tetanus or in contraction executes a series of minute successive movements, which the author calls *shocks*, which accumulate to produce muscular contraction, he has examined the discharge of the torpedo, and found there a similar complexity. Having passed this discharge through an electro-magnetic tracing apparatus he found that it was made up of minute shocks, which recur at the rate of 150 in a second. Cold reduces the rate alike of the muscular and electric shocks, whilst heat acts inversely. Hence the author concludes that these two functions are really homologues. The gymnotus gave similar results to those of the torpedo. The latter fish, when connected with a telephone and slightly excited, produced a very short croaking. If a prolonged discharge is occasioned by pricking the electric lobe of the brain the sound produced lasts three or four seconds and in tonality borders upon *mi* (165 vibrations).

Does the Didymium of Samarskite Differ from that of Cerite?—Lecoq de Boisbaudran.—The author concludes that both give alike the three blue rays 482.2, 475.8, and 469.1.

New Spectral Rays in Substances Extracted from Samarskite.—Lecoq de Boisbaudran.—On examining with the spectroscope both by absorption and by means of the electric spark, the products of his operations on the mixture of earths from samarskite, the author has observed rays or bands not to be referred to any element formerly known, and not corresponding to the descriptions of the spectra of the earths recently announced by MM. Delafontaine, L. Smith, Soret, and De Marignac. These new rays of absorption and emission seem to belong to one and the same body. The emission spectrum is composed of four bands shaded towards the left and formed of narrow rays, the strongest of which is the most refrangible and forms the right margin of the band. The absorption spectrum comprises two strong bands in the blue, and several rays of less importance in the green. The metal which yields these new spectra is precipitated as a double potassic sulphate along with didymium; its simple sulphate is rather less soluble than that of didymium; its oxalate is precipitated along with didymium, but ammonia separates the oxide of the new metal before that of didymium.

Unequal Propagation of Light Polarised Circularly in Bodies Submitted to the Action of Magnetism According to the Direction of Magnetisation and of the Luminous Vibrations.—H. Becquerel.—The phenomena of magnetic rotatory polarisation is accompanied, like natural rotatory polarisation, with a variation in the speed of propagation of two luminous rays polarised circularly in an inverse direction.

Compressibility of Gases at High Pressures.—E. H. Amagat.—Under a pressure of 430 atmospheres the volume of a gas is nearly $\frac{1}{2}$ greater than that which is deduced from the law of Mariotte.

Improvements in Harrison's Electric Lamp.—E. Ducretet.—This paper is not intelligible without the accompanying figure.

Relations which Connect the Tetric and Oxytetric Acids and their Homologues to Succinyl, Malyl, and other Radicles of the Bibasic Acids.—M. E. Demarçay.—The nature of the radicle of oxytetric acid is that of malyl, the radicle of malic acid.

Bromo-citraconic Acid.—E. Bourgoïn.—Not adapted for useful abstraction.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 8, February 20, 1879.

A Last Word on Matter.—E. Lafond.—“I announce to our friends that *matter is known*. I know now what it is. I will confine myself to telling you that it is essentially the *negation of being*! *Inanis et vacua*. This is a great step for science and the coronation of my system. We float in God, like the fish in water. The latter knows its element; man denies his. Which is the fool?”

Improvement in Bunsen's Battery.—F. Lefebure.—The oxidation of the exterior surface of the zincs contributes nothing to the liberation of electricity, so it may be safely covered with varnish, thus reducing the consumption of zinc by one half.

Poisonous Properties of Phenic Acid.—According to M. E. Küster the surgical use of this acid is not unattended with danger. He has collected twenty-six cases of evident poisoning, more or less severe. He recommends thymol, salicylic acid, or dilute solutions of zinc chloride (8 per cent) as antiseptic dressings.

An Unfortunate Pharmacist.—A French pharmacist has been fined 625 francs for illegally practising medicine. He had sold a few sous worth of “white water” (Goulard's lotion) for the treatment of a burn. The patient died of tetanus and the pharmacist was accused of poisoning, but finally escaped with the penalty above mentioned.

The Birth-rate in France.—Dr. Bertillon calculates that France capitalises $1\frac{1}{2}$ milliards yearly by reason of its small birth-rate, whilst Germany loses $1\frac{1}{2}$ milliards yearly by the rapid increase of its population.

Preservation of the Eggs of the Silk-worm in Different Gaseous Media.—G. Luvini.—The author has preserved the eggs of the silk-worm in hydrogen, carbonic acid, oxygen, and nitrogen for about three months. Those which had been kept in carbonic acid and in nitrogen after removal hatched well and gave an almost complete yield.

Verhandlungen des Vereins zur Beförderung des Gewerbfleisses. Part 1, January, 1879.

This part contains no chemico-technological matter.

MEETINGS FOR THE WEEK.

- MONDAY, 17th.**—Medical, 8.
— Society of Arts, 8. “Dwelling Houses Their Sanitary Construction and Arrangements,” by Dr. W. H. Corfield, M.A. (Cantor Lectures.)
— London Institution, 5.
TUESDAY, 18th.—Civil Engineers, 8.
— Royal Institution, 3. “Animal Development,” Prof. Schäfer.
— Zoological, 8.30.
— Society of Arts, 8. “Africa, a Paramount Necessity for the Future Prosperity of the Leading Industries of England,” by James Bradshaw.
WEDNESDAY, 19th.—Society of Arts, 8. “Economical Gardens for Londoners,” by W. Mattieu Williams, F.R.A.S., F.C.S.
— Meteorological, 7.
THURSDAY, 20th.—Royal, 8.30.
— Royal Institution, 5. “Sound,” Prof. Tyndall.
— Royal Society Club, 6.30.
— Chemical, 8. “On Perplumbic Ethide,” by E. Frankland and A. Lawrance. “On the Decomposition of Water by Certain Metalloids,” by C. F. Cross and A. Higgin. “On the Volumetric Determination of Chromium,” by W. J. Sole. “The Production of the Higher Oxides of Iron, Chromium, Manganese, and Bismuth,” by W. Foster.
— London Institution, 7.
— Zoological, 4.
FRIDAY, 21st.—Royal Institution, 9. “Denotating Agents,” by Prof. Abel.
SATURDAY, 22nd.—Royal Institution, 3. “Etching,” by Mr. Seymour Haden.
— Physical, 3. “On Selective Reflection,” by Capt. Abney, R.E., F.R.S. “On the Fracture of Colloids,” by Dr. F. Guthrie, F.R.S.

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 - VI. On Electrical Insulation in High Vacua. By William Crookes, F.R.S.
 - VII. Spider's Web for Micrometers.
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THE CHEMICAL NEWS.

Vol. XXXIX. No. 1008.

NEW PATENT LEGISLATION.

AMONG the many remedies proposed for the present depressed condition of British manufactures it is more than strange that such an immense majority overlook the simplest means—the encouragement of invention by giving increased facilities for obtaining and retaining patents. When we consider what we owe to invention in the past; when we further reflect that our most formidable manufacturing rivals possess a patent law in most points superior to our own, and that they ascribe to this patent law a very great share in their rapidly-growing industrial prosperity; when we reflect on these things, it does indeed seem to us surprising that patent law reform is not recognised as *the* question of the day.

Our present patent system, though possessing certain meritorious points, labours under very serious drawbacks. Its capital defect is the exorbitant cost of obtaining and upholding a patent, which amounts to £175 in stamp duties alone. It is very true that of this sum £50 are payable before the end of the third and £100 before the expiration of the seventh year respectively. But this plan of periodical taxes, non-payment of which at once renders the patent void, is fatal to the most essential element in the value of every kind of property, viz., security. It is not to be expected that manufacturers will pay a royalty to work a patent which if they only wait for a year may perhaps be thrown open to the world.

If we turn to the United States we find that there one payment of 35 dollars secures to an inventor the absolute right to his ideas for a term of seventeen years. Can we wonder if under such a system working men spend their leisure in trying to devise some improvement, some new "notion," of a useful character, and that foreign inventors are rapidly learning to consider America as the most favourable field for the practical development of their ideas?

Our national apathy on this most important subject is, however, not quite universal. Messrs. Anderson, Mundella, Dalrymple, and Alex. Brown are about to bring in a Bill which, should it become law, will be a very great improvement. The duration of patents is proposed to be extended from fourteen years—the present limit—to twenty-one. The stamp duties are to be reduced to £85, of which £10 are payable on obtaining the patent, £25 at the end of seven years, and £50 at the end of fourteen years. Patents already in force may be extended to twenty-one years, and shall be exempt from the payments formerly prescribed at the end of the third and of the seventh year, if not already due, and shall be liable merely to the payments prescribed in the new scale. These proposals, if carried out, would be a boon alike to inventors and to the nation at large. There is extension of time, reduction of cost, and though the vicious system of successive duties is not wholly abandoned, yet every patent, not otherwise voidable, is safe for seven years, which will give the patentee some chance at least of getting it adopted in practice.

Unfortunately, in opposition to this wise and moderate Bill, a measure is to be brought in under the auspices of Government which

"— keeps the word of promise to our ear
And breaks it to our hope."

The duration of a patent is to be extended to twenty-one years. But this boon is to be purchased at a cost of another £100, payable at the end of the *twelfth* year for the very suspicious reason alleged in the memorandum of explanations that the "public"—i.e., pirates—"may

have long notice of the patentee's intention to renew beyond the fourteen years." We almost fear that in these words lurks a possibility that the continuance of a patent for the remaining seven years may be opposed. The stamps duties payable on obtaining the patent are, indeed, to be reduced to £12 10s., but with the further duties at the third, seventh, and twelfth year the total cost will be increased to the enormous sum of £262 10s., whilst the opportunities lapsing from non-payment will be increased from two to three. As a still further blow to security of possession we find the following most mischievous clause:—"A patent shall be liable at any time after the end of three years from its date to be revoked on either of the following grounds:—(a) That the patentee fails to use or put in practice the invention to a reasonable extent within the United Kingdom, or to make reasonable efforts to secure the use or practice thereof there, proof to the contrary whereof shall lie on him; or (b) that he fails to grant licences to proper persons requesting the same, on terms which the Lord Chancellor deems reasonable."

When we remind our readers that in the general opinion of practical men it is decidedly exceptional for a patent to be brought into operation so early as the third year, except the inventor happens to be a capitalist, they will agree with us that this clause, if not in intention a cunningly-devised proviso for the confiscation of patent right, will act as such in practice. Men who would otherwise have come to terms with the patentee will, if this clause becomes law, wait till the third year has expired. We admit that a patentee who works his invention in some foreign country and refuses to do so in the United Kingdom might justly be held to have forfeited his rights. But this end could surely be attained without in substance enacting that no poor man shall hold a patent for more than three years.

To give the introducers their due, the Bill contains a few good features. The time of provisional protection before giving notice to proceed is extended to nine months, and if a stamp duty has not been paid at the proper date from accident or mistake the Lord Chancellor may, on petition, enlarge the time so as not to exceed three months longer. We know instances where patentees for want of this arrangement have been compelled to apply for a Private Act of Parliament—a most costly remedy. There are also increased facilities for amendment, disclaimer, &c., even after sealing.

Still, notwithstanding these minor improvements, if we consider that the cost of a patent, instead of being greatly reduced, will be augmented, and its character of permanence within the allotted term as greatly decreased, we can only pronounce this measure a step in the wrong direction, which it is the duty of all friends of the industrial progress of the nation to oppose by every means in their power.

ON VAPOUR-DENSITIES.

By JAMES T. BROWN.

Now that V. Meyer and C. Meyer have perfected a vapour-density method, which is not only extremely simple but is performed so rapidly that it will render previous methods matters of history, it seems to me that the only thing to prevent beginners from shirking the subject, as they certainly do, is to simplify the formulæ. The vapour-density must be—

wt. of substance

— wt. of air displaced by vapour

and when expressed in that form it is easily intelligible, but although different chemists find it convenient to arrange their formulæ in various ways, still when they publish their methods they should express them in the typical form. If (taking the weight of a c.c. of air at 0.0012932) instead of using Meyer's constant 587780 we

employ its reciprocal, we have the factor 0'00000170157, and then the formula becomes—

$$\frac{S}{(B-w)V \frac{0'00000170157}{1+0'00367t}}$$

As this factor is the weight of a c.c. of air at 0° C. and 1 m.m. pressure, it is the basis of the tables that I calculated. The very brief remarks that I made at the Chemical Society referred solely to the formulæ of the methods based on the *overflow* of mercury, and were an introduction to the hope that I expressed that someone would thoroughly investigate the very interesting subject of vapour-tensions at high temperatures.

186, Wandsworth Road, S.W.
March 5, 1879,

ON MANGANESE STEEL.

By SERGIUS KERN, M.E., St. Petersburg.

THE author had an opportunity of analysing some steel shavings from a broken pinion-wheel. It was stated that the pinion was in use for only a month, in the coupling of a rolling-mill for rails, when the pinion suddenly cracked in two halves; some of the pinion teeth were also damaged. The fracture had a very fine grain. Analysis gave the following results:—

	Middle Part. Per cent.	Broken Teeth. Per cent.
Sulphur.. .. .	0'10	0'09
Phosphorus	0'27	0'27
Copper	0'01	trace
Manganese	1'50	1'51
Carbon	0'65	0'65

These results prove the metal to be the so-called manganese steel. Certain metallurgists tell us that such a metal resists shocks very well, and that this enormous quantity of manganese added to the metal neutralises the evil effects of phosphorus. But the best thing that can be done is to avoid the use of such cast-metal cured by manganese. Two strips of the same metal when white-hot were welded together, but when it was attempted to bend them the first blow divided the strip into two pieces through the weld. Steel must not contain more than 0'3 to 0'4 per cent of manganese, and a metal containing 1'50 to 2'00 per cent of manganese is in most cases good for nothing. Inferior material, if even cured by manganese, will always give inferior steel. But as certain works prepare such a curious steel, it would be a benefit for the buyers if the manufacturers would supply them with a true analysis of the steel, because otherwise the buyer may get a cast-metal which only the manufacturer can call "steel."

THE MANUFACTURE OF POTASSIUM IODIDE.

By E. SCHERING.

THE following three methods are in actual use:—

- (1.) Decomposition of barium iodide (obtained from barium sulphide and iodine) with potassium sulphate.
- (2.) Introduction of iodine into caustic potassa; evaporation to dryness, and fusion with carbon in order to reduce iodate.
- (3.) Decomposition of ferroso-ferric iodide with potassium carbonate.

Satisfactory results can be obtained by any of these methods, and the choice must be determined by local considerations.

In method No. 1 the preparation of a barium sulphide

of high and regular strength is not unattended with nuisance, and the lixiviation of the barium sulphate requires much time. On the other hand, potassium sulphate can be obtained cheaper and purer than the corresponding carbonate, whilst the barium sulphate can be readily utilised for the reproduction of sulphide.

Method No. 2 obviates the necessity for washing a precipitate, and yields at once a very strong solution of potassium iodide; but the preparation of pure caustic potassa, and the concentration and subsequent fusion are circumstantial and tedious. The author therefore prefers the third method, as ferroso-ferric iodide is easily prepared and the carbonate of iron is readily washed.

To obtain cubic crystals of a porcelain-like appearance it is essential in the first method to ensure the complete decomposition of the barium sulphide by the iodine: if alkaline sulphides are mixed with the potassium iodide the crystals are paltry. If the lye contains iron sulphide, which is soluble in hot and concentrated potassium iodide, the crystals take a blue appearance. An excess of iodine dissolves foreign metals present in the barium sulphide, and the crystals may then be discoloured.

In the case of the second method, irregularity in melting may produce iodic acid, and a caustic potassa not free from sulphates causes the presence in the lye of alkaline sulphide. Both these injurious impurities must be removed prior to crystallising.

In the third method these annoyances are excluded. Salts of sodium must in all cases be avoided. Some manufacturers, to avoid the presence of sulphides, leave purposely a trace of iodate in the lye. The result is that the crystals turn yellow. The presence of lead in the iodine is exceedingly objectionable. This metal is soluble in concentrated potassium iodate, and cannot be precipitated by sulphuretted hydrogen except after great dilution.

If not removed lead affects not merely the colour but the forms of the crystals.

No demonstrable trace of potassium carbonate is admissible either for medical or photographic purposes. Potassium iodide, therefore, should be unaffected by salts of barium. The perfect absence of chlorine can never be attained, as even the best sample of iodine as well as of potassium carbonate contain traces of this impurity.

The Chilian iodine, obtained from soda-salt-petre, is becoming a formidable rival to the European product, which cannot be offered at a reduced figure as the manufacturers have lost their market for potassium chloride in consequence of the rivalry of the Staassfurt mines. Chilian iodine is met with in commerce either as a paste or as copper iodide.—*Chemische Industrie.*

CHEMICAL CALCULATIONS.

WE have received from Mr. Lupton, Professor of Chemistry at Harrow, a card destined to assist beginners in the solution of "chemical problems." By this expression the authors means, of course, not the many unsolved questions in chemical science with which so many highly disciplined minds are now grappling, but the problems which are given to pupils as means of instruction, and which, according to Mr. Lupton, "now form a large part of the chemical training at many schools." The card seems to us useful and well arranged for this purpose. It comprises a table of forty-one of the more abundant elementary bodies with their atomic weights and the corresponding logarithms. There is also a table of constants, chemical and physical, with their logarithms.

Whilst we have great pleasure in acknowledging the value of these tables, we cannot help considering it a misfortune that chemistry should be taught merely from books and calculations. We do not deny that the student may thus acquire an extensive and a fairly accurate knowledge of chemical facts and of chemical laws. But the

main educational value of chemistry, and, in like manner, of physics and biology, as it seems to us, lies herein—that these sciences, practically taught, train the mind and the outward senses to deal with objects, natural or artificial, and to draw conclusions from facts observed. They thus supply a grievous want in the ordinary system of English public school education where the intellect has been, till lately, solely engaged with words and with abstractions.

ON INDIGO-BLUE
FROM
POLYGONUM TINCTORIUM AND OTHER
PLANTS.

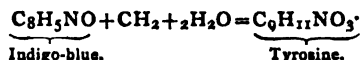
By EDWARD SCHUNCK, Ph.D., F.R.S.

SOME papers read before the Society many years ago, and subsequently published in its "Memoirs,"* contain an account of my experiments with the leaves of the *Isatis tinctoria*, or common woad, the well-known plant employed in Europe for dyeing blue before the introduction of indigo from the East. I showed that the leaves of this plant do not, as some have supposed, contain either indigo-blue or its hydride ready formed, but yield by careful treatment a peculiar glucoside—indican—which, when acted on by acids and other reagents, splits up into indigo-blue and indigo-glucine, the latter being a body resembling glucose. My experiments also show that this substance, indican, is a highly unstable body, undergoing when its watery solution is heated for some time, or, more rapidly, by the action of caustic alkalies, an entire change, on the completion of which it no longer yields indigo-blue by decomposition with acids, but in place of the latter gives indigo-red, indifulvine, leucine, and other products. Though I succeeded in ascertaining the composition of indican and the relation in which it stands to indigo-blue, the difficulty of obtaining large quantities of it in consequence of its excessive liability to change, prevented my proceeding further with the investigation. It seemed to me, however, that it might be of some interest to ascertain whether other indigo-yielding plants contain ready-formed indigo-blue (as has been maintained with so much persistence), or whether the colouring-matter exists in the vegetable cells in the form of indican or some other glucoside; and I have accordingly examined such of the plants known to give indigo as I have been able to procure.

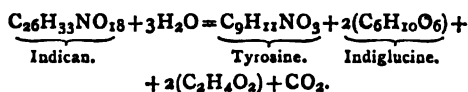
Before stating the results to which I wish to direct attention on this occasion, I may mention an observation belonging, strictly speaking, to the part of the subject previously treated of, which, however, I will now describe in a few words, as I may not have another opportunity of doing so. In my last memoir I stated that, among the products of decomposition of indican from *Isatis tinctoria*, leucine is usually found, sometimes, indeed, in considerable quantities. In some more recent experiments made with woad leaves I obtained, besides leucine, a substance having all the properties of tyrosine. This substance was only slightly soluble in cold water, but soluble in boiling water, from which it separated on cooling in long needles, forming, when dry, a snow-white felted mass. Its watery solution gave the well-known reaction with mercuric nitrate. Its solution in concentrated sulphuric acid, after neutralisation with barium carbonate, gave a purple colour with ferric chloride. According to Prof. Gamgee, who had the kindness to examine the substance for me, it showed under the microscope the forms characteristic of ordinary tyrosine. There could be no doubt therefore, of its identity with the latter. As the tyrosine in this case was not obtained from pure indican, but from the crude alcoholic extract of the leaves, it is impossible to say whether it pre-existed in the plant or whether it was a pro-

duct of decomposition of the indican of the extract. The latter supposition is the most probable one; for tyrosine being almost insoluble in alcohol could hardly be contained in any appreciable quantity in the alcoholic extract of the leaves.

Some connection between tyrosine and indigo-blue has frequently been suspected by chemists. Indeed a glance at the formulæ of the two bodies will show that some connection is possible, since, by replacing H by CH₃ in indigo-blue and adding 2H₂O, we arrived at the formula of tyrosine, thus:—



In order to explain the formation of tyrosine from indican we may suppose the latter to split up into tyrosine, indigluceine, acetic acid, and carbonate dioxide, thus:—



Supposing the tyrosine in this case to have been really formed from indican, the question suggests itself whether the leucine and tyrosine so frequently found in the animal organism as products of disease, may not be derived from some substance similar to indican rather than directly and immediately from albuminoids.

Polygonum tinctorium.

This plant has long been known and employed as a source of indigo by the Chinese. According to Stanislas Julien*, who has given translations from Chinese works of various process for extracting indigo from the leaves, the plant is called in China Lán, the most productive variety being termed Tcha-Lán, i.e. the Lán resembling the tea shrub. It was introduced into Europe in the eighteenth century, and at one time, particularly about the years 1838 to 1840, formed the subject of numerous investigations by eminent French botanists and chemists, such as Turpin, Joly, Baudrimont, Pelletier, Robiquet, and others, some hopes being entertained that the plant might be cultivated profitably in France. The numerous trials made with this view having led to no result, the matter fell again into oblivion, and this interesting plant remained what it was before, a mere curiosity.

I obtained the seeds of the plants from Messrs. Vilmorin, Andrieux, and Co., the eminent horticulturists of Paris, and therefore felt sure of their genuineness. They were sown in a hotbed, and germinated rapidly. As soon as the young plants were a few inches high they were transplanted into the open ground, where they grew vigorously, producing an abundance of leaves and attaining during the summer season a height of nearly three feet. Towards the end of summer spikes of pretty pink flowers, resembling those of other species of *Polygonum*, made their appearance. The seed, however, did not ripen in the open air, the plant being cut down by the early frosts before this could take place; but, by growing a few plants under glass, I obtained a quantity of well-matured seeds, which yielded another crop in the following season.

For a botanical description of *Polygonum tinctorium* I must refer to Turpin† and Joly,‡ the latter of whom has given a full account of its structure and affinities. Of the various organs, the leaves, being the seat of the blue colouring-matter, are alone of any interest to the chemist. These leaves, which are large, oval in shape, and glossy, show no indication of the presence within their tissue of any pigment besides the chlorophyll to which they owe their lively green colour, except, occasionally, in certain places where they have suffered injury from the bites of

* *Comptes Rendus*, t. vii., p. 703.

† *Comptes Rendus*, t. vii., p. 806.

‡ *Sur le Polygonum tinctorium*: Montpellier, 1839.

* "Memoirs," ser. 2, vol. xii., p. 177, and vol. xiv., p. 181.

insects or from others causes, and where blue spots make their appearance—a phenomenon which I shall endeavour to explain presently. It is nevertheless easy to show that they contain a considerable quantity of what the French call a *matière colorable*, i.e., a substance which, though colourless in itself, yields colouring-matter by appropriate treatment. A few leaves having been cut into pieces and rubbed up with a little water in a mortar to a thin paste, the mass is poured on a bit of calico, and yields, by squeezing and kneading, a green muddy liquid which, on the addition of a little sugar of lead solution, gives a green flocculent precipitate containing the chlorophyll, albumen, and other matters previously held in suspension. The liquid filtered from this precipitate is clear and yellow, and on being mixed with sulphuric or hydrochloric acid and left to stand for several hours, yields a deposit consisting of tolerably pure indigo-blue. The amount of colouring-matter obtained in this way from *Polygonum tinctorium* is far greater than that which the same quantity of wood-leaves grown in the same locality would produce, proving that the yield is influenced not only by soil and climate, but also by the peculiar nature of the plant.

The isolation of the *matière colorable* of *Polygonum tinctorium* is still not a very easy task. The same precautions must be observed as in the case of *Isatis tinctoria*, particularly as regards the evaporation of the solvents employed, which must always be effected without applying artificial heat. Unless some means are at disposal for evaporating rapidly at the ordinary temperature, by means of a current of air or otherwise, success is very uncertain. The method formerly employed in preparing indican from *Isatis tinctoria* was first tried. The leaves of the plant were dried in a stove moderately heated, and, while still warm, ground to powder. The powder, which had the colour of fresh hay, was passed through a hair sieve to separate the leaf-stalks and other fibrous portions, and then extracted in a percolator with spirit of wine. The green alcoholic extract was evaporated in a shallow tin dish, the evaporation being assisted by passing a current of air over the surface of the liquid in the apparatus formerly described. Chlorophyll and fatty matter were deposited during evaporation, leaving a brown watery liquid, which was poured off, agitated with freshly precipitated copper oxide, and filtered. The copper in the filtrate was precipitated with sulphuretted hydrogen, and the filtered liquid was evaporated as before. The residue was treated with absolute ether, which dissolved a portion, and left, on evaporation, a yellow syrup. This syrup is the indigo-producing body as pure as it is possible to obtain it. I prefer to this the following process, as being more expeditious and surer. The alcoholic extract of the dried leaves having been evaporated, the watery liquid which is left is mixed with acetate of lead solution, which gives a dirty yellow precipitate, consisting of chlorophyll and other impurities in combination with lead. To the clear yellow filtrate basic lead acetate is added; this gives a primrose-yellow precipitate, which is filtered off and, after being washed with water and then with alcohol, is suspended in absolute alcohol, through which a current of carbonic anhydride is passed. After the gas has passed through for some time the liquid acquires a yellow colour, and, after being filtered from the insoluble portion, consisting principally of lead carbonate, is evaporated at the ordinary temperature. Water added to the residue leaves a portion undissolved, which is filtered off. Sulphuretted hydrogen is passed through the filtrate to precipitate the lead contained in it; and having been again filtered, it is evaporated, when it leaves a syrupy residue which may be treated with ether as before.

The indigo-producing body thus obtained is, if it be permitted to draw a certain conclusion from mere qualitative reactions, identical with the indican of *Isatis tinctoria*. Its appearance is that of a yellow transparent syrup, showing no tendency to assume a crystalline form. It is soluble in water, alcohol, and ether. The watery solution has a more or less acid reaction. It becomes of a deep yellow

colour on the addition of caustic alkali, and gives with basic lead acetate a light yellow precipitate. When the watery solution is mixed with a little sulphuric or hydrochloric acid and left to stand for some time, the surface of the liquid becomes covered with a film of indigo-blue, a deposit of the same substance being usually formed at the bottom. The filtered liquid shows, when tested with a salt of copper and an excess of caustic alkali, the well-known reaction of glucose. If, however, the watery solution is left to stand at the ordinary temperature for a considerable time, or if it is simply boiled for some time, or if it is mixed with caustic alkali and left for a short time, it no longer yields indigo-blue on the addition of acid. This is probably due, as I have shown to be the case with indican, to a molecular change, resulting, when completed, in the formation of a body which, when decomposed with acids, yields indirubine and brown resinous substances in place of indigo-blue. When a large quantity of watery solution is mixed with acid and left to stand, a portion of the substances undergoes, it seems, the same change; for the deposit formed when operating with one litre or more of the solution contains not only indigo-blue, but also indirubine, indifulvine, and other products. The deposit, which in this case is almost black, after being filtered off, washed, and dried, is treated first with caustic alkali and then with cold alcohol, in order to remove the indifulvine and other resinous substances. On treating the residue with boiling alcohol, the indirubine dissolves, and is obtained, after several crystallisations, in the beautiful dark-red needles characteristic of the substance. The portion insoluble in boiling alcohol is indigo-blue, requiring for its purification merely to be dissolved in some suitable menstruum, such as boiling aniline. I am inclined to think that the same molecular change takes place in the cells of the plant during the later stages of its development; for I obtained from some leaves gathered late in the season when the flowers had begun to appear, a quantity of indican having the usual appearance, but giving, by decomposition with acid, far less indigo-blue and more indirubine and other products than the indican from younger leaves. This result was confirmed by experiments, to be described presently, made with the leaves themselves.

(To be continued.)

NOTICES OF BOOKS.

A Practical Treatise on the Manufacture of Sulphuric Acid. By A. G. Lock and C. G. Lock. London: Sampson Low, Marston, Searle, and Rivington.

We have here a thoroughgoing and a practical book on what may fairly be pronounced the most important branch of industrial chemistry. The authors, unlike too many compilers of so-called technological works, do not content themselves with generalities, but enter closely into working details. The construction of the kilns and chambers, with their accessory arrangements, the selection of materials, the points needful for economy in management, and the prevention of waste and nuisance, are all carefully and clearly presented to the reader, with the important aid of seventy-seven "construction plates" drawn to scale and showing the chief recent improvements.

It may seem strange, if not somewhat humiliating, that in a manufacture so old and so widespread as that of sulphuric acid, and depending so little on manipulative niceties, there should be so much divergence of opinion concerning arrangements of plant and methods of working. Still, from another point of view, it may be held consolatory to know that one, and that not the smallest, of our national industries is capable of very decided improvements if those who combine the opportunity and the requisite knowledge will give the question the careful examination it deserves.

Our authors begin with a consideration of the raw

material, a subject which derives additional importance from the growing objection to the industrial uses of arsenical products, and from arsenic no pyrites are absolutely free. It remains to be seen whether the vast deposits of sulphur in Iceland may not enable us to overcome this difficulty, situate as they are so much nearer our shores than are the Sicilian mines. We notice that Messrs. Lock quote from Mr. H. A. Smith the statement that in certain Irish pyrites the arsenic present exceeds 2 per cent. They are also, on an average, low in sulphur, and are very hard, if not altogether impossible, to burn reasonably clean—a fact which goes far to vitiate the details of production-cost quoted from Messrs. Richardson and Watts as to the relative economy of brimstone, of Belgian, and of Irish pyrites. In a subsequent part of the work the authors express a preference for the best-class Norwegian pyrites beyond all others.

The observations on the analysis of sulphur ores are too brief to be of much practical value, and might have been advantageously omitted, especially as this subject is discussed at length in other works.

The structure of kilns for brimstone and for different qualities of pyrites is treated very fully, with the aid of plates representing the special arrangements for a rich Norwegian ore, for a poor Irish, and for the poor English sulphur ores from the coal-fields, commonly known as "coal-brasses" or brass lumps. There are, besides, representations of the kiln fronts of Messrs. Daglish and Co., and of Gerstenhofer's kiln, now very extensively, and we believe successfully, used on the Continent.

The capacity, shape, and arrangement of chambers are considered in the second chapter. Here, it must be owned, practical men are very far from having come to a unanimous conclusion as to the best construction—an evident proof that we have here much still to learn. The authors quote with disapprobation, or at least with strong doubt, the dimensions recommended by Mr. H. A. Smith, viz.—"Length, 160, 200, or even 210 feet; width, 30 feet, increasing in ratio to length; and height, 10 feet, not to be increased for any length of chamber." As grounds for rejecting these proportions they allege the great consumption of lead in proportion to bulk, and they show that a chamber 210 feet by 30 feet, having only 1 inch of acid all over the bottom—the very smallest depth needful to prevent the escape of the gases—will lock up 23 tons of acid. As the opposite extreme they mention Hasenclever's proportions, where the minimum height of the chambers is fixed at 32½ feet to 130 feet in length and 32½ feet in breadth. It surely ought to be possible to decide which of these two glaringly opposed systems gives in practice the best results, and whether either of them is preferable to the less extreme dimensions recommended by the authors, viz.—in a set of 60 feet long, 27 wide, and 23½ feet high, and 30 feet long, 23 feet wide, and 22 feet high.

The proposal to inject "atomised" or pulverised water into the chambers in place of steam is next passed under review, as also the suggestion to admit a solution of nitre in the form of spray. But on these questions the authors, though citing the opinions of others, do not appear to have formed any decided conclusion of their own.

The evidence adduced on the proposal to use free nitric acid instead of nitrate of soda seems decidedly favourable to this system, which is almost universally adopted on the Continent. In some German works the consumption of nitre in this form has been brought as low as 1½, 1¼, and even, it is said, 1 per cent of the sulphur present in the pyrites. The nature of this economy will be fully appreciated if we remember that where no Gay-Lussac's or Glover's tower is in use some manufacturers use 9 or even 10 lbs. of nitre to every 100 lbs. of sulphur actually burnt, and that even where such contrivances are at work the proportion often reaches 5 per cent. Dr. Lunge, whom we consider an excellent authority, considers the free acid so much superior to the nitre that nothing but a fear of having too much difficulty with the workmen in the first instance would deter him from discarding the latter system in England.

This last remark suggests some unpleasant reflections. One of the many reasons which place us at a disadvantage in competing with foreigners is the obstinacy with which the British workman clings to traditional methods, and his disinclination to give fair play to any new invention which he may be called upon to adopt.

The chapter on "chamber construction" is admirably complete and practical.

In treating of "condensers and columns" the evidence for and against the use of the Glover tower is fairly stated. The authors consider it certain that in large works, at least, this tower will become as common an adjunct as the Gay-Lussac column.

The succeeding chapters on the concentration of the acid, on the utilisation of waste products, and on accessories must be pronounced satisfactory, though we regret that want of space forbids us to examine them in detail. The final sections on attempts to manufacture sulphuric acid without the use of chambers, and on the so-called Nordhausen or fuming acid, will also be found valuable and suggestive. The latter acid is now in extensive demand in the arts, e.g., in preparing the sulpho-conjugated acids of certain coal-tar colours, and in the purification of ozokerite. As shippers and railway companies dislike undertaking its conveyance there is the greater need for its production on the spot.

In concluding this necessarily brief sketch, we beg to record our opinion that though no reading will enable a man to dispense with actual experience, yet that the authors have done all that can be done in the form of a book. Every chemical manufacturer, every scientific chemist who is at all connected with practical operations, has room for gratitude to the authors.

The book is admirably got up, and reflects great credit on the publishers.

Our Domestic Poisons, or the Poisonous Effects of Certain Dyes and Colours Used in Domestic Fabrics. By H. CARR, M.Inst.C.E. London: Ridgway.

ARSENIC is in more senses than one a delicate thing to handle. On a somewhat similar occasion we ventured to put to the accuser a few questions, such as would have been urged by counsel if the matter had come before a court of law, and in consequence we were half buried beneath a heap of indignant letters. In the present pamphlet it is maintained that "national health is suffering from the use of arsenic and other poisons in the manufacture of domestic fabrics," and the author takes in hand to arouse public feeling with the end of demanding legislative interference. For this purpose he has put himself in communication with certain eminent physicians and chemists, and has obtained from them evidence more or less confirmatory. The first point which here strikes us is that Mr. Carr has not—as far as here appears—applied to any chemist specially and practically acquainted with the arts of dyeing and calico-printing and able to inform him what poisonous colours are actually in use and what are the circumstances of their application. The instances of poisoning given, at least, as far as textile fabrics are concerned, are by no means as explicit as could be wished. We seek to know in such cases the name and address of patient, medical adviser, and chemist, if one has been called in; the date of the occurrence, and the precise name of the colour said to have occasioned the mischief. The terms "mauve-dyed articles," "beautiful red, scarlet, and mauve colours," &c., are too vague to be employed in so serious a question. These demands may perhaps seem stringent, but it would be difficult to name any subject which has given rise to so many newspaper canards as the alleged poisonous character of dyes. Our contemporary, Dr. Reimann, of the *Färber Zeitung*, has been at the pains of sifting not a few of these stories. In many cases it was found on rigid scrutiny that the alleged sufferer and his doctor were both non-existent. At other times the source of the disease has been declared to be

some dye which is not in the market at all, or which, if known, could not by any means produce the colour in question. The celebrated shoe-maker of Stettin, whose inflamed eyes were first ascribed to an arsenical hat-lining, but were afterwards traced to a diligent internal use of potato-whiskey is a case in point. We have no objection to Mr. Carr collecting instances of poisoning from dyes, but they should be such as will bear cross-examination. As far as arsenic is concerned we must bear in mind that there are many channels other than dyes and pigments through which it may be introduced into the human system. If we remember the violet powder case we may well suspect that arsenic is one ingredient of the cosmetic powders which so many ladies apply to their hands and faces.

As regards the aniline colours their physiological action is still a matter of doubt. That aniline itself is poisonous is far from being a proof against its derivatives. Much research will therefore be needed before either physicians or chemists can form a decided opinion on the safety of any given dye or pigment, and such research, thanks to the Vivisection Act, has been rendered in England all but impossible. Meantime we should do nothing rashly. If we burden our colour makers, dyers, and printers with regulations from which their competitors in France, Germany, &c., are free, we shall merely strike a heavy blow at our national industry. In none of those countries is there any restriction on the use of non-arsenical coal-tar colours in textile manufactures.

The British Journal Photographic Almanac and Photographer's Daily Companion for 1879. Edited by J. TRAILL TAYLOR. Henry Greenwood.

THE twenty-fifth issue of this well-known year-book gives a vivid idea of the immense progress which photography has made in this country since first it put forth its few modest leaves. Besides the usual information contained in almanacs we have 200 pages containing succinct accounts of every photographic process and discovery belonging to the past year, condensed from an immense number of scientific *Transactions* and journals from all parts of the world, prefaced by a summary of the progress made in photography during the past year. Mr. Taylor secedes from the editorship of the Almanac this year. May we suggest to his successor the advisability of classifying the information given in the body of the work under distinct headings instead of allowing it to be mixed up in picturesque disorder as at present. There is a copious index appended to the book. As an instance of the enterprise of photographic material dealers we may mention that the almanac contains nearly one hundred and fifty octavo pages of advertisements.

The Patentees' Manual: being a Treatise on the Law and Practice of Letters Patent, especially intended for the Use of Patentees and Inventors. By JAMES JOHNSON, Barrister-at-Law, and J. HENRY JOHNSON, A.I.C.E., Solicitor. Fourth Edition. Longmans and Co. 1879.

This is the fourth edition of Messrs. Johnson's standard work, and is a great advance on the preceding ones. The whole text has been thoroughly revised, and several of the chapters greatly enlarged or re-written. A new chapter on Oppositions to the Grant of Patents has been added, and the latest decisions of the Courts incorporated with the text. The new German and Spanish Patent Laws have been given, and the Appendix contains an account of those of our Colonies, as well as a reprint of all the Acts of Parliament bearing on the subject, with references to typical cases decided thereon. Messrs. Johnson may be congratulated on the successful way in which they have respectively completed their arduous task. Their work is another instance of the truth of Adam Smith's grand theory of the division of labour. There is a copious index at the end.

CORRESPONDENCE.

TENACITY OF STARCH.

To the Editor of the Chemical News.

SIR,—I have read with interest the method employed by Mr. George Whewell for determining the tenacity of starch, but think it right to call attention to the last paragraph of his paper, which gives the conclusions he draws from his experiments; these in my opinion are entirely erroneous. He says in effect, If the paste given by one sample of starch is firmer or has more "tenacity" than that given by another, the former will give more stiffness to the cloth into which it is introduced in direct proportion to the resistance which it offers to a weight placed on its surface. If this be so, then supposing we take calcined farina (British gum), and dissolve it by boiling in the smallest proportion of water possible; it would make a syrup which would not support any weight placed on its surface, and consequently, according to Mr. Whewell's theory, would not give any stiffness to cloth when used in sizing. And yet it is undoubtedly the case that if we make solutions by boiling equal weights of British gum and farina with equal weights of water, and pass threads or cloth through each, and then dry them, the former solution will give more stiffness to the threads or cloth than the latter, and weight for weight the farina which possesses the *least* "tenacity" would produce the greatest stiffness, because the thinner the sample boils the more easily will the solid matter be able to enter the fibre, and it will enter in greater quantity. I know that the opinion expressed by Mr. Whewell is held by many manufacturers, but it is fallacious.—I am, &c.,

WILLIAM THOMSON.

Royal Institution, Manchester.

CHLORIDE OF CALCIUM.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxix. p. 97, Mr. O. Gluge, of Sarrebrück, in a "Note upon Chloride of Calcium," refers to its manufacture as a by-product in the manufacture of alkali by the Solvay process; but he does not seem to be aware that it is also a by-product in the manufacture of bicarbonate of soda by the Leblanc process and of bleaching-powder by the Weldon process. We have made chloride of calcium for sale by one or both of these processes for upwards of fifteen years, during a considerable portion of which time our sales have exceeded thirty tons weekly. If Mr. Gluge will refer to the back numbers of the CHEMICAL NEWS he will find that for many years our product was advertised in your pages.—We are, &c.,

GASKELL, DEACON, AND CO.

THE INSTITUTE OF CHEMISTRY.

To the Editor of the Chemical News.

SIR,—The letter of Mr. Charles E. Groves, written in answer to mine signed "A Disgusted Promoter," commences by accusing me of inaccuracy, the only instance adduced being that I referred to the recent conference as one on the "Adulteration of Food and Drugs" instead of the "Adulteration of Articles of Food." It is true that Mr. Groves also points out that when my letter appeared the conference which I spoke of as pending had already taken place, but you, Sir, if my original MS. is still extant, will know that my letter was *written and received* by you a few days *before* the date fixed for the second conference. There is another error in my letter which I will give Mr. Groves the benefit of. I spoke of the conference on trade

certificates being held in December, whereas the true date of the meeting was November 22.

Although it was true when I wrote my last letter that no report of the November conference had been published, the statement is no longer correct, for *since my letter appeared in your pages* the members have received a printed report of the proceedings. Will Mr. Groves excuse the suggestion that its circulation is a *consequence* of my criticisms? Whatever may be the cause, it is satisfactory to find that the Council have published the report, though it has taken them *nearly four months* to do it.

I may remind your readers that I did *not*, as stated by Mr. Groves, "recommend Fellows not to pay their subscriptions," but merely suggested that they should defer doing so until the Council proposed to spend the large sum of money already in hand. Mr. Groves will best serve the Institute by publishing a candid statement of the plans of the Council. To gain this was the object of my last letter, but Mr. Groves wholly ignores my challenge, in his reply, though from the fact that the Report of the Council has just been circulated amongst the Members I conclude that he is alive to the nature of it.

My letter having secured the chief results aimed at, I beg to subscribe myself

AN APPEASED PROMOTER.

March 15, 1879.

VICTOR MEYER'S NEW METHOD OF VAPOUR DENSITY DETERMINATION.

To the Editor of the Chemical News.

SIR,—Referring to the interesting note by Greville Williams, F.R.S., in the CHEMICAL NEWS (vol. xxxix., p. 110), I see he mentions having abolished the funnel, *d*, figuring in the woodcut (page 66). Now this funnel-like aperture is somewhat exaggerated in the printing, and in reality the opening or widening is very little larger than the main stem of the instrument, and just conveniently sufficient to admit a small india-rubber stopper, which V. Meyer always uses. The advantage of the transparent glass vessel or bath, *c*, is that the ascent of the heated vapours can easily be seen on boiling the liquid—the bulb—and so the operation is more conveniently regulated. I may also mention that Prof. Victor Meyer has not only had the instrument made of porcelain (better for several reasons than metal), but has made and is making by means of this instrument (identical in every other respect with the glass one) a number of interesting determinations of the vapour-densities of certain extremely high boiling inorganic substances, leading to still more interesting theoretical conclusions.—I am, &c.,

WATSON SMITH, F.C.S., F.I.C.

Zürich, March 17, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 8, February 24, 1879.

Induced Currents Resulting from the Movements of a Coil across an Electro-Magnetic System.—M. Th. du Moncel.—The induction currents in action in the Gramme machine consist of those which result from the movement of the covered spirals before the inducer and of those determined by the inversions of the polarity of the iron ring.

Observations with Reference to M. G. Planté's Recent Work "Researches on Electricity."—E. Becquerel.—A secondary battery of 800 elements arranged

on M. Planté's system, which may be easily charged by means of two nitric acid elements, can give effects of tension equal to those of 1200 nitric acid elements. These currents are of course temporary, and the apparatus serves a kind of condenser of voltaic currents, but they are sufficiently permanent to produce mechanical, thermic, and luminous effects of great power.

Hemihedral Forms of Alums.—Lecoq de Boisbaudran.—The author presented to the Academy a crystal of alum having four smooth and four rugged surfaces, arranged so as to represent the union of two inverse tetrahedra. It was obtained by steeping an octahedron of chromo-potassic alum for some hours in a slightly super-saturated solution of basic ammonia-alum.

Resistance to a Change of Condition of Crystalline Surfaces in Presence of their Mother-liquor.—Lecoq de Boisbaudran.—The action of different isomorphous crystals upon the solution of any one of them is not the same. The transition from the state of the very slow dissolution of a crystalline surface to its very slow increase is not effected abruptly in consequence of an extremely slight change in the concentration of the liquid, but each surface remains intact without either gaining or losing substance in a mother-liquor, the strength of which ranges within certain limits, which, though narrow, are easily perceived. The resistance to a change of state is modified for each system of surfaces independently, in such a manner that an alteration in external conditions (a change in the composition or in the temperature of the liquid, &c.) involves generally a change in the relation of the resistances of the two given systems of surfaces. Contrary to the opinion of some authorities, there does not exist a state of mobile equilibrium between a crystalline surface and its mother-liquor; there is no continual interchange of molecules, but merely a continual erosion or a continual deposition, and within the limits of resistance to change of condition there is neither erosion nor deposition.

Reflections on the Communication made by M. de Lesseps Concerning the Contagion of the Plague.—M. Bouillaud.—The author considers that the researches of M. Pasteur on the low organisms considered as propagators of disease must greatly modify our views on the spread of epidemics.

Projection of Molecular Shadows.—W. Crookes.

Action of Various Coloured Light upon a Layer of Silver Bromide Saturated with Various Organic Colouring Matters.—Ch. Cros.—Upon plates sensitised with an extract of mallows the direct spectrum of the Drummond light is inactive in the medium green, but very active at the red and the violet extremities. With carthamin the mean portion is most active, and with chlorophyll the action extends throughout the visible spectrum, and even somewhat beyond. It presents several distinct maxima and minima.

The Production of Crystalline Barium Chromate.

—L. Bourgeois.—The author heats to bright redness for half an hour 2 equivs. barium chloride with 1 equiv. potassium chromate and 1 equiv. sodium chromate; lets the mass cool slowly, when crystals of a pistachio-green are found in its interior. They are freed from the alkaline chlorides by washing with boiling water. The specific gravity is 4.60. They dissolve readily in dilute hydrochloric or nitric acid, giving an orange liquid. They are decomposed by sulphuric acid with formation of barium sulphate and chromic acid. They consist of 60.4 per cent of baryta, and 39.6 of chromic acid, corresponding to the formula BaO, CrO₃. The green colour is not due to the presence of chromic oxide.

Composition of Beer-yeast.—P. Schützenberger and A. Destrem.—The authors consider yeast as containing complex compounds, at once hydrocarbonic and proteic, formed like glucosides, and easily decomposed by acids and alkalis. The exterior of the granules differs from

the interior merely by containing more hydrocarbonic matter.

Pyrogenous Carbides of American Petroleum.—L. Prunier.—The author has discovered a compound of the composition $(C_{12}H_2)_n$, and containing 97.29 of carbon. These figures indicate the existence of a new group of incomplete carbides higher than any previously known, since benzerythren, parachrysen, fluoranthren, and pyren do not contain more than 95 per cent of carbon.

On Glycide.—M. Hanriot.—Glycide represents the alcohol corresponding to epichlorhydrin.

Production of Aniline-black by Means of the Chromates in Presence of Chlorates.—S. Grawitz.—The author denies the accuracy of the conclusions of M. Witz on the inutility of chromates.

*Verhandlungen des Vereins zur Beforderung des
Gewerbfleisses.* Heft 10, December, 1878.

This number is chiefly taken up with a paper on the theory of "Closed Air Machines." There are official communications concerning boiler explosions and the storing of petroleum.

La Correspondance Scientifique.

No. 36, February 18, 1879.

Three veins of coal have been discovered in the immediate neighbourhood of Lake Nyanza. One of them is 7 feet in thickness, and the others respectively 3 feet and 1 foot.

According to *Le Petit Marseillais* the waters of the Dead Sea are about to be worked for potash, iodine, and bromine. The projectors calculate on delivering potassium chloride in London at 90 francs per ton, thus underselling the Stassfurt manufacturers.

Chemiker Zeitung.

No. 1, 1879.

Preparation of Zinc from Blende.—The blende is partly roasted and is then treated in a muffle with burnt lime and coke. Calcium sulphide is formed and the greater part of the zinc present in the ore is volatilised. This is condensed or collected as zinc oxide as in the ordinary zinc furnaces. From the calcium sulphide sulphuretted hydrogen is evolved, and by its action upon the sulphurous acid gas given off on roasting the blende sulphur is obtained.

Process for Obtaining Tanning Agents along with Cellulose, Gum, and Acetic Acid from Vegetable Matter.—Dr. A. Mitscherlich.—Wood or other vegetable matter is treated at an elevated temperature with bisulphite of lime (under pressure?). The cellulose which is cemented together by other compounds is set at liberty in the same state in which it exists in plants. It is readily freed by washing from the accompanying substances, and can be at once utilised for paper-making, &c. The soluble substances vary according to the material employed, and embrace compounds suitable for tanning and for the manufactures of gum, acetic acid, and alcohol. The methods for separating these substances are omitted.

Dr. H. Buff, Professor of Physics at the University of Giessen, died on December 24, 1878.

Sawyer and Man's electric lamp has been tried in New York with great success. The light is produced in an atmosphere of nitrogen so that there can be no combustion and no generation of nitrogen oxides—an objection lately raised against the electric light. The cost for equal illumination is said to be only one-fortieth that of gas, and the light can be regulated from a faint gleam up to the intensity of 30 gas-jets.

Analysis of Butter.—E. Mylius finds that butter globules if examined with the microscope by polarised

light can readily be distinguished from other melted fats, as the latter display a crystalline structure.

Quality of Milk.—According to the researches of Berguerel, Vernois, and Müller the respective proportions of the constituents of milk are affected not by race but by age, diet, and other causes not yet ascertained.

No. 2, January 9, 1879.

Quantitative Spectrum Analysis.—Wolff proposes the following methods for the determination of cobalt, copper, and metallic iron. 1. Determination of very small quantities of cobalt.—A solution of cobalt-sulphocyanide, obtained by mixing a very dilute alcoholic solution of chloride of cobalt or cobaltous nitrate with an excess of alcoholic ammonium sulphocyanide, displays in the region $C_{39}D-C_{55}D$ a very characteristic absorption spectrum. By means of a solution of cobalt sulphocyanide of known strength he determines the residual strength of light in a stratum of 1 centimetre in thickness, the coefficient of extinction and consequently the absorptive proportion for cobalt. The latter number is multiplied by the volume and the coefficient of extinction of the solution under examination, and gives the weight of the cobalt present. It must be remarked that in these experiments alcohol of the exact specific gravity 0.833 must be used, because an alcoholic solution of cobalt sulphocyanide differs in the intensity of its blue colour according to the strength of the alcohol. This behaviour has been utilised by Morrel for a colorimetric determination of alcohol. 2. Determination of Copper.—The absorptive proportion for copper sulphate or for metallic copper is measured as ammonio-sulphate of copper. To determine small quantities of copper in articles of food, &c., the ash is dissolved in an acid, the dissolved copper is separated electrolytically, re-dissolved in nitric acid, and saturated with ammonia to a certain volume. The latter multiplied by the coefficient of extinction of the solution obtained and with the absorptive proportion of metallic copper as previously ascertained, shows the weight of the copper present in solution. 3. Determination of Metallic Iron.—The ferri-ferrous substance is mixed with a normal cupric solution, whose volume, coefficient of extinction, and equivalent in iron are known. The solution filtered from the deposit of copper is evaporated, oxidised, super-saturated with ammonia, separated by filtration from ferric oxide, and made up to a fixed volume. The coefficient of extinction and the equivalent in iron of this solution are ascertained; the latter subtracted from the known iron equivalent of the normal copper solution originally employed shows the metallic iron.—*Zeitschrift Anal. Chem.*, 18, 38.

Simplified Butter Test on Hehner's Principle.—Reichert saponifies a known quantity of butter, dissolves the soap in water, adds sulphuric acid, and distils. If the butter is pure 2½ grms. of the sample yield a distillate which requires at least 13 c.c. of decinormal soda.

MISCELLANEOUS.

Professor Church and the Agricultural College.—At a meeting of the Committee of Management of the Agricultural College, held last week in London, present Earl Bathurst, the Earl of Ducie, Col. Kingscote, C.B., M.P., Mr. G. Sotherton Estcourt, M.P., Mr. A. L. Goddard, M.P., Mr. T. S. Bazley, and Mr. Edward Bowly, the following resolution was passed, in reference to Prof. Church's application for permission to reside out of the College:—"The Committee of Management are of opinion that the discipline of the Agricultural College cannot be satisfactorily maintained except by the residence of professors within the College walls in conformity with the original by-law No. 47. Being fully sensible of the services rendered by Prof. Church during his 16 years' residence in the

College, they the more regret that they cannot accede to his recent proposal of non-residence, a compliance with it involving such alterations as would unduly disturb the present organisation of the College staff." We understand that, in consequence of the above decision as to Prof. Church's connection with the Agricultural College, the two resident professors next to him in seniority have resigned their respective chairs; the chair of mathematics and physics being vacated by Prof. H. W. Lloyd Tanner, M.A., that of natural history by Prof. Fream, B.Sc., F.G.S.

Science in the Streets.—Under the title of the "New Parisian Electric Pipe-lights" a number of itinerant vendors are selling in the streets of the metropolis small brass boxes containing three or four rodlets of metallic sodium, an inch long and one-tenth of an inch square. The method of using these lights, which are sold at a penny a box, is simple. A morsel of the metal is pinched off and placed on a piece of paper which has first been moistened with water or saliva. The sodium of course inflames and sets alight to the paper. The risk to person and property that will be caused by the indiscriminate distribution of such a very dangerous material as metallic sodium, which even skilled chemists are obliged to handle with the greatest care, is obvious to all who know its properties. We have on a former occasion called attention to the evil of allowing the open sale of metallic sodium in toy shops under the name of "Satan's Tears," and we believe a stop was put to it by the police. In the present instance the danger is much greater, inasmuch as any urchin who has a penny to spare has it in his power to set half a street in flames at a moment's notice.

NOTES AND QUERIES.

Gases from Vitriol Chambers.—Would any of your readers describe and explain an easy method of testing the exit gases from vitriol chambers, and state what is considered "good working." I may say we have not Gay-Lussac or Glover towers, but simply one packed with coke as a condenser.—W. L.

Substitute for Litmus.—Where can I procure small quantities of Porrier's "orange 3," "orange 4," or tropæolin OO, recently recommended as a substitute for litmus by Mr. Greville Williams?—VOLUMETRIC ANALYST.

MEETINGS FOR THE WEEK.

- MONDAY, 24th.**—Medical, 8.30.
— Society of Arts, 8. "Dwelling Houses: Their Sanitary Construction and Arrangements," by Dr. W. H. Corfield, M.A. (Cantor Lectures.)
— London Institution, 5.
— Royal Geographical, 8.30.
TUESDAY, 25th.—Civil Engineers, 8.
— Royal Institution, 3. "Animal Development," Prof. Schäfer.
— Anthropological, 8.
WEDNESDAY, 26th.—Society of Arts, 8. "The Treatment of Iron to Prevent Corrosion," Prof. Barff, M.A.
— Geological, 8.
THURSDAY, 27th.—Royal, 8.30.
— Royal Institution, 3. "Sound," Prof. Tyndall.
— Philosophical Club, 6.30.
— London Institution, 7.
— Society of Arts, 8. "The Inoxidation of Iron and the Coating of Metals and other Surfaces with Platinum, by the Processes of M. Dode," L. M. Stoffel, C.E.
FRIDAY, 28th.—Royal Institution, 9. "Geography of the Oxus, and its Changes," Sir Henry C. Rawlinson.
— Society of Arts, 8. "The Practicability and Advantage of a Ship Canal Through the Island of Rami-eran," Simon McBean, C.E.
SATURDAY, 29th.—Royal Institution, 3. "Etching," by Mr. Seymour Haden.

TO CORRESPONDENTS.

W. H. D.—We do not recollect.
C. W. W.—The best way would be to advertise it.
More Light.—Either pronunciation of the word appears to be equally correct.

The Prestolee Alkali Works, Farnworth, near Bolton, Lancashire, fitted with costly Plant, Machinery, and Apparatus for the Manufacture of Soda-ash, Bleaching-powder and Liquor, and Sulphuric Acid, in complete working order, with possession.—Preliminary.

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THE CHEMICAL NEWS.

VOL. XXXIX. No. 1009

THE ROYAL AGRICULTURAL COLLEGE,
CIRENCESTER.

PROFESSOR CHURCH'S connection with the Royal Agricultural College will, we regret to say, shortly be severed, the only reason being, as far as we can ascertain, that he is about to marry, and therefore will be unable to reside in the College. Seeing, however, that his immediate predecessor was non-resident, we should have thought the Council would have hesitated to adopt this as their reason for depriving the College of the services of so able and experienced a chemist as Professor Church, and we shall look with some eagerness for their explanation of the circumstances.

It appears that at the beginning of February Prof. Church, who has occupied the Chair of Chemistry at the College for the last sixteen years, had an interview with the Principal, the Reverend John Constable, and informed him of his intention to marry during the coming vacation, and of henceforth discontinuing to reside in the College. This intimation the Principal regarded as a resignation of the Professorship, and wrote to Professor Church accepting the resignation. In answer to this letter, Professor Church informed the Principal that if he decided that his Chair became vacant by the fact of non-residence it would therefore become vacant at the date previously mentioned, April 24th. He furthermore assumed that this decision was that of the Governing Body of the College, as he was elected in 1863 under the bye-law then in force, which placed the power of appointing and dismissing Professors in the hands of that body, and not under the twenty-fourth bye-law of 1870 (*vide* CHEMICAL NEWS, vol. xxxix., p. 105), by which this power was vested in the Principal. The matter was then referred to the Committee of Management, and on the 5th of March the following resolution was passed:—

"The Committee of Management are of opinion that the discipline of the Agricultural College cannot be satisfactorily maintained except by the residence of the Professors within the walls of the College in conformity with the original bye-law No. 47. Being fully sensible of the services rendered by Professor Church during his sixteen years' residence in the College, they the more regret that they cannot accede to his recent proposal of non-residence, a compliance with it involving such alterations as would unduly disturb the present organisation of the College staff."

This decision has not only deprived the College of the services of Professor Church, but of those of his two senior colleagues, Professors Lloyd Tanner and Pream, who have respectively vacated the Chairs of Mathematics and Physics and Natural History as a protest against Professor Church's unworthy treatment.

The bye-laws of the College which we published in the CHEMICAL NEWS are perfectly unique in their way, and would seem to show that the Principal entirely supersedes the Committee of Management. He evidently considers that the Students of the College are a set of unruly schoolboys who are only to be kept in order by the strictest discipline, and that the Professors must lead the life of monks. They are not only to devote themselves day and night to the service of the College, but they are to act the much-despised part of the "*pion*," or *maitre d'étude* of a French *Lyce*, and see that their young charges do not tear their clothes, tell fibs, or say naughty words. But as three Professors would have still been left to reside in the College

after Professor Church had left, and as the College has frequently been left in the charge of two, the argument that Professor Church's proposal for non-residence would unduly disturb the present organisation of the College carries with it very little weight.

The consequences of the resignation of three senior Professors, the only University men, by the way, in the College, cannot be otherwise than disastrous. At the end of this term the senior Professor will be a young man, who only two years ago was a student himself, and he will very possibly be called on to rule over students who are his seniors in years, an anomaly which cannot fail to bring about unpleasant consequences sooner or later.

We hear that a new bye-law has just been promulgated for the well-being of the future Professors. They are not to engage in any literary work without special leave from the Principal; in other words, they must take no steps to make a reputation outside the College walls without the express permission of their ruler. Had this rule obtained during Professor Church's Professorship how many valuable contributions to scientific literature would have been lost.

Such restrictions will certainly prevent any man of worth from accepting a position in the College. The Principal may, it is true, gather round him a knot of second or third rate men; but the interests of the College and, indeed, of agriculture generally, must necessarily be injured thereby; indeed, we do not see how the College can escape collapse and ultimate extinction if something is not done either by the Government or the bondholders to prevent such a catastrophe.

On looking over the list of Professors that have formerly occupied Chairs in the College we find that Principal Constable has made no less than twenty-two within the last fifteen years, more by many, we should imagine, than any other man in a similar position has made in his whole life. Professor Church has remained at the College for sixteen long years, but most of the others have left but little trace behind them. Again and again have first class men been engaged, but there has never been any inducement held out to them to cast in their lot with the College.

A contemporary gives the Principal the credit of having made the College pay, the bondholders having of late received a four per cent dividend, and something has been done towards paying off the loan, but this has only been accomplished by raising the fees from £90 to £126, by discontinuing the farm which some authorities consider to be a breach of the provisions of the Charter, the sale of the steam plough, and the most pernicious economy in every direction. Nothing is spent on the Museum, and nothing in several departments for specimens and apparatus. To have a good surplus every year is no doubt gratifying to the Council and bondholders, but even gold may be purchased at too high a rate.

We feel sure, in conclusion, that our readers will join us in expressing our sincere sympathy with Professor Church and his colleagues at thus having to sever themselves from an institution with which they were so long connected, and to which they had rendered such valuable services, as well as in paying a just tribute of admiration at the manly, straightforward, and honourable course they have taken.

Progress Effected in the Preparation of Colours Derived from Coal.—Adolphe Kopp.—The attempts made to dispense with arsenic in the manufacture of magenta have only been partially successful. The process of Coupier, who causes nitro-benzol to react upon a mixture of aniline and toluidin in presence of hydrochloric acid, is adopted in some manufactories, but it offers serious difficulties which interfere with its general employment. It is more costly and more difficult manage.—*Moniteur Scientifique*.

FILTER-PRESSES FOR CHEMICAL WORKS.

By J. MARZELL.

In this time of depression of trade it is natural that chemical manufacturers should pay more attention to those kinds of apparatus which, by a surer and better working, enable them to increase the yield, and permit quicker passing through the intermediate stages of the processes, thus giving to a plant a higher working power and efficiency. Among that class of machinery the so-called "filter-presses" undoubtedly hold one of the foremost places, and their being taken up so rapidly—especially in the last year—surely speaks in their favour.

The principle of the filter-presses is generally known; it is a filtering under pressure, for the object of—

- (a) Filtering the largest possible quantity of fluid—consequently solids—in comparison—through the smallest possible filter space.
- (b) Freeing the residue remaining in the filter as much as possible from the filtrate.

Both points are realised by the pressure applied in filling the filter-presses, and generally amounting to 120 lbs. per square inch. At the same time the filter-press represents the most convenient arrangement of filter cloth, as a very large surface is brought into a very small space. In the largest size presses over 425 sq. feet effective filtering surface are arranged in a box, if I may call it so, of $4\frac{1}{2}$ sq. ft. to 9 sq. ft. General rules of yields cannot be laid down, as specific gravity and physical properties of the different stuffs are important points to deal with. The greatest difficulty for chemical works always was to find a press suiting the different stuffs and operations, as these in most cases cannot be altered or modified without losing on the other side all advantages the filter-presses might have promised. It was therefore only natural that the massive and enormous wooden cubes brought out at first, with their elaborate handling, had soon to make room for lighter constructions, with a number of improvements in many directions.

An important improvement has recently been made by M. A. L. G. Dehne. I refer to a new arrangement for an absolute outwashing of the press cakes in the press itself under the following conditions:—

- (a) To use as little wash water as possible.
- (b) To remove the last traces of soluble substance out of the cake, to wash absolutely or completely—be it either to wash the cakes, as the valuable part "neutral," as I may term it (as required with alizarin, tartrate of lime, blanc fix, &c.), or to separate the valuable soluble substances from the worthless residues without any loss, and to get the wash liquors in a concentrated state, the latter being of great importance for a following concentration for crystallisation (as with sugar, citric, tartaric acids, alum, sulpho-salts, &c.)
- (c) To do all that in the shortest time possible.

In the old method of washing the water was introduced on the top of the cakes, trickling through the same, taking up in its way the soluble substances it met with, and leaving the press by the cocks. Every manufacturer acquainted with these presses will have found that it takes a long time, and is seldom possible, to remove the last parts of soluble stuff, and therefore it was preferred in many cases to take out the cakes, boil them up with water, and to re-filter-press them; in many cases even repeating the operation several times.

The reason of this improper acting of the washing apparatus is the insufficient distribution of the wash liquor in the cakes, caused—

- (a) By the construction.
- (b) And principally, by the air enclosed in the cakes.

Especially is this last point of great importance. That the amount of air in the cakes is consider-

able is proved by the fact that in the manufacture of china and faience the cakes of clay coming out of the presses never can be used on the chairs before having passed cutting and kneading machinery, as in baking the air would form bubbles on the goods. In the washing process the air prevents the wash liquor from a thorough penetrating of the cakes. The water coming from above would not remove but only dislodge the air. The consequence of that was the conclusion to let the wash liquor enter at the bottom and make its way upwards, driving thus the air before it, which is removed by a system of air channels and escapes, with a whistling noise, out of a mouthpiece.

The other disadvantage of the old system was the following:—The water entered on the upper edge of the cake in two streams of about $\frac{1}{2}$ in. thick. By the pressure the washing was applied and the two streams naturally tried to find the most direct way to the outlet, and that is the line drawn from their point of entrance to the outlet cocks! Besides that, the way was regulated by the little air-bladders forming a regular kind of network in the cake. The result was that the streams, instead of equally impregnating the material, only parted themselves in the cakes, so to say, forming a system of small rivers, the islands of which never came under the influence of the water.

To do away with this irregularity M. Dehne covered the channelled surfaces of the press plates inside; the chambers with perforated sheets of metal, and now in washing brought the water into the space formed between the channelled surface and the perforated plate. Thus on both sides of the cakes in this way walls or plates of water are produced, which by the hydraulic acting pressure are driven diametrically into the cakes. With a cake of 1 in. thick the water has only to move $\frac{1}{2}$ in. to join the liquor coming from the other side. As the air has been removed beforehand, each molecule of the stuff in the presses is now brought into contact with the water, which thereby has full power to take up all soluble particles. The arrangement allows at the same time to regulate the pressure under which the extraction takes place.

So far the theory. The practice, by working on large scales under different circumstances and with different materials, has given the best results.

As example I give the returns of the K. K. Zuckerfabrik Swolonowas (Austria), which were kindly placed at my disposal. The material is the product of precipitating saccharate of lime with carbonic acid, thus yielding carbonate of lime, mixed with sugar syrup, which latter has to be removed.

"The presses are such with 18 chambers of M. Dehne's, showing 125 square ft. filtering surface proved for 120 lbs. per square inch."

Filling of press = 30 minutes; washing absolutely = 18 minutes; emptying and starting = 6 minutes; together 54 minutes.

The weight of the extracted cakes out of one press, 315 lbs. (157 k.); the cakes are hard.

The extracted cakes were *entirely free from sugar*.

The quantity of wash-water used for one press = 39 gals. (176 litres). Density of the entire wash-liquor = 5.23° balling and 4.11 per cent sugar.

The filter cloth has not to be removed for a fortnight. (This for the reason in the arrangement that the wash-liquor takes the opposite way of the filtration, thus always cleaning the pores of the cloth.—J. M.) The firm receives daily 13½ tons (13,627 k.) of the washed-out residues. The corresponding residues of the old presses retained about 2.46 per cent of sugar, which it was not economical to regain, and the firm reckon the *net gain* they have by putting up Dehne's new presses with absolute extraction with 47½ tons (47,400 k.) of sugar in a campaign of 150 days, representing a value of £1422 (14,220 Austrian florins).

The quicker working and the advantage of getting pretty

concentrated liquors into the evaporating pans need not be mentioned.

Another example. The quick extraction of valuable cakes is given in the manufacture of the artificial alizarin material; alizarin paste must be freed from the sulphate of soda. With the old presses the washing was going on twelve to fourteen hours before the charge of a press with 18 chambers was finished. The new presses with absolute extraction—according to the returns of an alizarin works using twelve of these new presses—only wanted 35 to 45 minutes for the absolute neutralisation of a properly precipitated paste. In this case a secondary but important advantage of the new process can be seen. In washing there is always some alizarin taken up into suspension and washed away, especially in the last hours, when the water is running pretty clear. As with the new presses the time, as well as the quantity of water, are decreased considerably, this loss is reduced to a minimum. Of great weight this point is in the treatment of such substances, which themselves are slightly soluble in water. I only mention tartrate of lime.

Notwithstanding their short life the new presses with absolute extraction have been taken up by a great many firms. Like most new things they have often to meet with mistrust and criticism in sometimes the most incomprehensible form. From the representatives of progress, however, "the go ahead people," the new apparatus everywhere meets with the greatest interest and acknowledgment of its practical advantages, and the large number of these presses already delivered to various branches of different countries may be proof of their value.

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ON INDIGO-BLUE FROM POLYGONUM TINCTORIUM AND OTHER PLANTS.

By EDWARD SCHUNCK, Ph.D., F.R.S.

(Continued from p. 120).

THE preceding experiments lead to the conclusion that the leaves of *Polygonum tinctorium* contain a substance not to be distinguished from the indican of *Isatis tinctoria*, which by decomposition with acids, yields indigo-blue and glucose, accompanied by some by-products, and that there is no proof of the existence of ready-formed colouring-matter in the plant while the latter is living and in a healthy state. The pre-existence of indigo-blue, or of its hydride, indigo-white, in these plants was taken for granted forty or fifty years ago when the class of bodies which we now call glucosides and the peculiar kind of decomposition which they undergo were unknown. Even now a superficial examination of some phenomena would almost certainly lead to the conclusion that the indigo-blue is formed by the action of air, i.e. in consequence of the oxidation of some easily oxidisable substance in the plant. Bearing in mind, however, with what extreme facility indican is decomposed, its watery solution on standing some time, even at the ordinary temperature, depositing indigo-blue, I think it will not be difficult to explain all the phenomena hitherto observed by myself and others.

On taking a plant of *Polygonum tinctorium* and making incisions with a penknife in the leaves between the main vessels, or crushing the soft parts of the leaves here and there with an agate pestle, then, after a short time plunging the whole plant into boiling alcohol to remove the chlorophyll, it will be found that those parts of the leaf which have not been injured, become white or retain only a faint yellow tinge, while those parts that have been cut, crushed, or otherwise injured, show a blue colour, the colouration extending for some distance inwards from the place where the lesion occurred, the most intense colour

being at the edge of the cut or bruise. So, too, in the living plant, when some injury accidentally occurs to a leaf, the part injured will appear blue. Nothing can be more natural than to suppose that in these cases the blue colouring-matter is formed by the action of the air, i.e. by the oxidation of some substance which escapes from the cells in consequence of organic lesion, just as the surface of a freshly cut apple or pear becomes brown on exposure.

If a plant of *Polygonum tinctorium* be immersed in water, and the water be frozen by surrounding the vessel containing it with a freezing-mixture, it will be found, after complete thawing, that the leaves or parts of leaves which have been thoroughly frozen appear of a dark colour and are quite flaccid; and if the plant be then immersed in boiling alcohol so as to dissolve the chlorophyll and other matters, those very parts show afterwards an intense blue colour, while those portions which had remained unfrozen appear almost colourless. This experiment, which had already been made by Joly, was considered by him to prove the pre-existence of indigo-blue in the plant—though why, if this were the case, the colouring-matter should not make its appearance in the unfrozen portions of the leaf, I am at a loss to understand.

The fact that a fresh leaf of *Polygonum tinctorium*, if immersed in alcohol or ether, appears blue after the chlorophyll has been removed, has also been considered to prove the pre-existence of indigo-blue in the cells. This phenomenon is always observed when the leaves are immersed in cold alcohol, and more distinctly when ordinary spirits of wine are taken than with absolute alcohol. A very simple experiment suffices, however, to prove that in this, as in all the other cases, appearances are deceptive. If freshly gathered leaves of *Polygonum tinctorium* are plunged at once not into cold, but into boiling alcohol, the whole of the colour is soon removed, the leaves retaining only a faint yellow tinge. On now simply evaporating the green alcoholic liquid, not a trace of indigo-blue will be found in the residue. It is therefore absolutely certain that the leaves contain no ready-formed colouring-matter; for so stable a body as indigo-blue could not possibly be decomposed or be made to disappear by the action of boiling alcohol only. It must necessarily appear either in the alcoholic extract of the leaves or in the residual portion left by the alcohol.

A very simple explanation offers itself, I think for all the phenomena hitherto observed. Indican, the mother substance of indigo-blue, is a body the molecules of which are in a state of unstable equilibrium. As long as it is contained within the cells of the plant the vitality of the cells keeps it in its original unchanged condition. As soon, however, as this vitality is destroyed (whether by organic lesion, by extreme cold, or by any other means), the indican begins to undergo decomposition, the molecules rearrange themselves in the order to which their chemical affinities predispose them, and the compound splits up into indigo-blue and indigluceine, this taking place so rapidly that in certain cases it would appear as if the colouring-matter pre-existed in the plant. If (to take the simplest case) the leaves as soon as gathered are immersed in cold spirits of wine or in cold ether, the vitality of the cells is thereby destroyed; and the indican contained in them is then in part decomposed, yielding indigo-blue, which remains undissolved, imparting a more or less distinct blue tint to the leaves. When boiling spirit of wine is taken, the indican is extracted before it can undergo decomposition, and dissolves in the spirit. It may be detected in the residue obtained on spontaneous evaporation of the alcoholic extract by its property of yielding indigo-blue on decomposition with acids as above described. It is possible that the leaves contain some ferment which hastens the decomposition of the indican as soon as vitality has ceased; but I have no positive evidence to offer in favour of this view.

I may, in conclusion, describe another experiment, which, though it teaches nothing new, confirms what I have just stated, and is interesting in its way. Having cut some

sprigs of *Polygonum tinctorium*, about six inches long, I immersed the cut ends in dilute hydrochloric acid (consisting of one part of acid of specific gravity 1.15 and ten parts of water), and left them to stand for several days exposed to the sun and air. The acid was gradually absorbed, ascending through the stems, first into the lower leaves, then into the higher ones. The gradual absorption of the acid was distinctly seen by the discolouration of the leaves, which commenced at the basis of each leaf and extended towards the apex, the lively green colour being changed into a dirty yellow. After some time this colouration was followed by a dark blue one, commencing at the base of each leaf and extending towards the apex, but never quite reaching the latter except in the lower leaves. When the change in colour had begun to show itself in the upper tenderer leaves, the whole showed symptoms of fading, all further power of absorbing the acid liquid seemed to be lost, and the sprigs were then at once immersed in hot spirit of wine. After remaining in the spirit until the chlorophyll was removed, the part of each leaf which had undergone a change appeared blue, whilst the part into which the acid had not penetrated appeared almost colourless. In this case it is probable that the indican was decomposed not so much in consequence of the loss of vitality in the cells as by the direct action of the acid. In some of the leaves there was another distinct blue colouration towards the apex, in the part to which the acid had not penetrated, separated from the blue part at the base by a white zone. This second colouration may be attributed to the loss of vitality in that part of the leaf.

All these experiments must be made with plants whilst in a state vigorous growth. If made when the season is advanced (that is, after the flowers have begun to appear), the leaves, though apparently unchanged, show only traces of blue colour after treatment and subsequent immersion in hot alcohol. This shows either that the indican has disappeared, being applied to other purposes in the economy of the plant, or that it has undergone the peculiar molecular change before referred to, into a substance which no longer yields indigo-blue by decomposition, but indirubine and other products. The latter is the more probable way of accounting for the difference; for the parts of the leaf which have been accidentally injured by the bites of insects or from other causes at the later stage of the plant's development become red, not blue, as at the earlier stages. This red colour disappears on immersion of the leaves in hot alcohol, the indirubine, to which it is probably due, being more soluble in that menstruum than indigo-blue.

The leaves of *Polygonum tinctorium* in which the blue colour has been developed by any of the means described, exhibit even to the naked eye, and still more distinctly when examined under the microscope, certain appearances which are not without some interest.

1. The colouring-matter seems to be confined to the parenchyma of the leaf. The stem and its fibrous ramifications in the leaf are free from it, so that in the coloured leaf the vessels may be distinctly traced as white veins on a blue ground. Even the cells of the parenchyma adjacent to the vessels are much less coloured than those a little further off, which produces the effect of a gradual shading of colour from the white of the vessels to the dark blue of the remoter cells. The cells of the leaf-cuticle are also free from colouring-matter.

2. The younger leaves at the summit of each branch generally show a more intense colour than the older ones near the base. Each leaf probably contains the same amount of colouring-matter; but in the lower leaves it is more widely distributed.

3. The intense and apparently uniform colouration of some of the leaves might lead to the conclusion that the cellular tissue is itself dyed blue—which would not seem improbable considering the affinity which indigo-blue shows for cellulose, as seen in the blue-dyeing of cotton fabrics. On examining the leaf-cells under the microscope,

this is found, however, not to be the case. The colouring-matter is discovered within the cells of the parenchyma in the shape of separate dots and parcels of various sizes, and apparently in an amorphous state, the cell-wall being quite colourless. These dots and parcels being very numerous, produce, when seen in the mass, a uniform blue colouration, more or less intense. The darker colour of some leaves is simply the effect of a greater crowding of the blue particles in each individual cell, the cells of the paler leaves containing fewer of these little masses, sometimes hardly any.

Mr. Charles Bailey, to whom I gave some specimens of leaves of *Polygonum tinctorium* coloured blue, had the kindness, at my request, to submit them to microscopic examination, and gave the following as his opinion thereon:—

"The colouring-matter left in these specimens would seem to be what Nägeli terms 'crystalloids;' and, with one exception, these bodies are, as far as my examination has gone, confined to the interior of the cells of the parenchyma. I do not see the least trace of any of this colouring-matter occurring in the intercellular spaces. The only part of the tissue where I find it, other than the parenchyma, is in the cells of the stomata; but it occurs nowhere else in the cuticle."

(To be continued.)

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, March 20, 1877.

Dr. J. H. GLADSTONE, F.R.S., President, in the Chair.

AFTER the announcement of visitors the minutes of the previous meeting were read and confirmed. The following certificates were read for the first time:—W. Johnstone, W. B. Turner, C. Slater, T. Palmer, A. H. Black, W. A. Bradbury.

THE PRESIDENT then called on Dr. FRANKLAND to read a paper on "*Plumbic Tetrethide*," by E. FRANKLAND and A. LAWRENCE. This substance was discovered by Buckton in 1859. The authors prepared the compound as follows:—Plumbic chloride is added to zinc ethyl contained in a stout glass bottle as long as any reaction takes place. The product is mixed slowly with a large volume of water, and distilled (from an oil bath) in a current of steam. The distillate separates into water and a heavy layer of plumbic tetrethide. During the preparation no gas was evolved, and the authors infer that the decomposition has two stages, $PbCl_2 + 2ZnEt_2 = PbEt_2 + 2ZnEtCl$ and $2PbEt_2 = Pb + PbEt_4$. The following gases have no action at ordinary temperatures on plumbic tetrethide:—Ammonia, carbonic anhydride, carbonic oxide, cyanogen, nitric oxide, oxygen, sulphuretted hydrogen. Sulphurous anhydride is rapidly absorbed and the liquid plumbic tetrethide converted into a white amorphous solid. This product was placed in a beaker, over which was inverted a second beaker, the two being luted with gummed paper. On heating the lower beaker with steam, crystals sublimed, which, after purification from a volatile lead compound by treatment with strong nitric acid, evaporation to dryness, and re-crystallisation from alcohol, proved in chemical composition and physical properties to be identical with diethyl-sulphone, SO_2Et_2 . The residue after the sublimation of the diethyl-sulphone consisted principally of plumbic ethyl-sulphinate,



Ethyl-sulphinic acid may be considered as sulphurous

acid, SOHO_2 , in which a semimolecule of ethyl has replaced one of hydroxyl, SOEtHO , or as propionic acid in which one atom of tetrad-sulphur has been substituted for one of carbon—



No volatile organo-metallic base was detected in the product of the action of SO_2 on PbEt_4 similar to that produced in the corresponding reaction with stannic ethide.

In answer to a question of the PRESIDENT,

Dr. FRANKLAND said that plumbic tetrathide could be prepared directly without the intervention of zinc ethyl, but the product was small in quantity and impure.

Prof. W. FOSTER then gave a verbal communication "On the Production of the Higher Oxides of Iron, Chromium, Manganese, and Bismuth." Prof. Foster said that the observations he had to make were a continuation of a paper which he had recently communicated to the Society. When a strongly alkaline solution of sodic hypobromite is heated with a solution of potassic ferrocyanide the solution rapidly becomes deep red from the formation of a ferrate. Almost any iron compound, and even freshly precipitated ferric hydrate can be substituted for the ferro-cyanide in the above reaction. The author has not attempted to isolate the potassium or sodium ferrate formed under these circumstances. Similar reactions occur if manganese, chromium, or bismuth compounds be boiled with alkaline hypobromite, permanganate, chromate, &c., being formed. Oxygen is evolved during the reaction. Cobalt, nickel, and copper salts also evolve oxygen. The author demonstrated this fact in the case of cupric sulphate, a considerable quantity of oxygen being collected when the sulphate was mixed with alkaline hypobromite, and the mixture heated to about 20°C .

Dr. WRIGHT said that some samples of bleaching-powder when closely stoppered were apt to explode: (he had always found manganese in these samples). Possibly this might be accounted for by the evolution of oxygen by a reaction similar to that described by the author.

Mr. NELSON asked if Prof. Foster had noticed any formation of a green manganate in the oxidation of manganese compounds. Some time since he was working with ferrates, and obtained once a green iron acid which he had never seen since. Perhaps by these reactions it might be obtained.

Mr. HARTLEY had noticed that when the copper sulphate and alkaline hypobromite were first mixed the solution turned yellow, but when the evolution of oxygen ceased the liquid was quite dark. This yellow substance was probably Brodie's cupric peroxide.

Mr. KIMOSSETT had made some experiments on the action of oxide of manganese on bleaching-powder; oxygen was evolved, but the gas was not pure.

Prof. FOSTER, in answer to a question of the President, said that he had not tested the oxygen evolved quantitatively as to its purity.

The next paper was read by the SECRETARY, "On the Decomposition of Water by certain Metalloids," by C. F. CROSS and A. HIGGINS. The statements of chemists with respect to the action of sulphur on water are conflicting. Mulder affirms that the vapour of water reacts on sulphur at a high temperature to form penta-thionic acid. Myers asserts that hydrogen sulphide and thio-sulphuric acid are formed. Girard also found that hydrogen sulphide was evolved by boiling sulphur with water. Geis ascribes this evolution to impurities in the sulphur. The authors therefore investigated the reaction of sulphur with water. Water containing "flowers of sulphur" in suspension was boiled, and the vapour conducted into lead acetate. The latter was continuously decomposed with the formation of sulphide. The quantity formed under various conditions was estimated. The reaction which occurs is probably $2\text{H}_2\text{O} + \text{S}_2 = 2\text{H}_2\text{S} + \text{SO}_2$. If air be present sulphuric acid is also formed. Some experiments

were also made with sealed tubes. The authors conclude that sulphur decomposes water, uniting both with its oxygen and hydrogen, the decomposition being independent of atmospheric oxygen. The authors repeated their experiments with sulphur purified by treatment with potassium permanganate, and find that the reactions above described remain unaltered. Selenium and tellurium have an appreciable action on water at 160° . Amorphous phosphorus does not decompose water at 100° . At 160° a small quantity went into solution. When lead acetate solution was substituted for water, metallic lead, phosphate, and phosphide were formed. With copper sulphate metallic copper was deposited with phosphide and sulphide, phosphoric and sulphuric acids being also formed; cupric chloride at 160° is first reduced to cuprous chloride, and finally converted into a phosphide. Vitreous phosphorus does not decompose boiling water except in the presence of oxygen; it reduces metallic (lead acetate) solutions even when oxygen is excluded. Bromine and iodine, when heated with excess of water, dissolve in small quantity as alkaline bromides, bromates, &c. If bromine be treated with lead acetate, lead bromide and dioxide are formed.

Mr. WARINGTON remarked that an Italian chemist had observed that gypsum was formed by the action of calcium carbonate on sulphur in the presence of moisture.

The next paper was read by the SECRETARY, "On the Volumetric Determination of Chromium," by W. J. SELL. The author having discovered that chromium in ordinary chromic salts can be completely converted into chromic acid by means of potassic permanganate, endeavoured to apply this reaction directly for the estimation of chromium. He was, however, unsuccessful, owing to the separation of manganese dioxide. He has devised the following method, which yields rapid and accurate results. The solution, containing chromium acidified with sulphuric acid, is boiled, and a dilute solution of permanganate added to the boiling liquid until a purplish tint remains after boiling for three minutes. The solution is then rendered slightly alkaline with sodic carbonate, alcohol is added, and the manganese filtered off. The chromic acid in the filtrate is estimated by titration with iodine and sodic thio-sulphate. The author has successfully applied the method to the estimation of chromium in chrome iron ore. He recommends the following plan of effecting its decomposition. The chrome iron ore is placed on the top of about ten times its weight of a mixture composed of one molecule of well-fused and powdered sodium bisulphate to two molecules of sodium fluoride, and the whole is ignited for fifteen minutes. An amount of sodium bisulphate is now added equal to that of the mixture taken, and when thoroughly fused a further addition of an equal quantity of bisulphate is made, the mass fused, and then rapidly cooled. The fused mass so obtained dissolves completely in boiling water acidified with sulphuric acid. In this way a determination can be made in an hour and a quarter. The author's attention has been directed to a notice by Mr. Wanklyn in the *Phil. Mag.*, February, as to the conversion of chromic oxide into chromate by alkaline permanganate. This result the author arrived at some months ago, but has not yet succeeded in applying it quantitatively.

During the reading of the last paper Dr. RUSSELL occupied the Chair.

The anniversary meeting will take place on Monday, March 31. The next ordinary meeting will be on April 3, when the following papers will be read:—"On Terpin and Terpinol," by Dr. Tilden; "On the Transformation of Aurin into Trimethyl-para-rosanilin," by R. S. Dale and C. Schorlemmer; "On a Gold Nugget from South America," by Mr. Attwood; "On the Solution of Aluminium Hydrate by Ammonia, and a Physical Isomeride of Alumina," by C. F. Cross.

PHYSICAL SOCIETY.

Ordinary Meeting, March 22, 1879.

Prof. W. G. ADAMS, President, in the Chair.

New member—Capt. Hastings R. Lees, R.N.

Capt. ABNEY, R.E., F.R.S., read a paper "*On Obtaining Photographic Records of Absorption Spectra.*" Absorption spectra have hitherto been recorded by the difficult process of hand copying; but the discovery by Capt. Abney of a silver salt sensitive to all rays in different degrees renders the photographic method available. The records thus obtained are photographs of the spectrum of the naked light of the source, and of that of the same light reduced by insertion of the absorbing material in its track, and these are taken parallel, so that the dark absorption lines can be readily compared. Examples of these were thrown on the screen. This method can be used as a new weapon in attacking solar physics, and determining whether or not compound bodies exist in the sun. Absorption spectra to compare with the sun's can be got for compound bodies by burning the matter in question in a flame in front of the slit, and passing a bright light through the flame.

Prof. GUTHRIE, F.R.S., then read a paper "*On the Fracture of Colloids.*" as illustrated by experiments on the breakage of glass plates either by pressure or heating at the centre or round the circumference. Circular plates of glass pressed at centre or circumference break in radial lines. However supported a plate breaks in the same fashion if heated in the same way. If heated in the middle the crack is peak-shaped, like an obelisk on a double pedestal, two cracks forming the outline, with sometimes a third down the middle. The two cracks unite before they reach the edge on one side, and (as afterwards pointed out by Prof. W. G. Adams) the three extremities of the two cracks all meet the edge at right angles to it. The crackage varies with the size and shape of the plates, the flame, and kind of glass; but the type is the same for all. Cracks cross each other. Prof. Guthrie defined a crack as the line where the ratio of cohesion to strain is least, and likened it to the lightning flash.

Mr. W. CHANDLER ROBERTS, F.R.S., said that he had observed once a volute spiral crack in dried hydrated silicic acid, and recommended Prof. Guthrie to study cracks in agate, which is the most perfect colloid known.

NEWCASTLE CHEMICAL SOCIETY.

Mr. J. W. SWAN in the Chair.

THE minutes of the previous meeting were read and confirmed.

Mr. John Cliff was elected a member.

The following names were read for the first time:—Mr. James C. Rollin, 1, St. Nicholas' Buildings, Newcastle-upon-Tyne; Mr. Thomas Douglas, Blaydon Manure Works, Blaydon-on-Tyne; Mr. Alfred Poole, Bottle Works, Blaydon-on-Tyne; Mr. W. F. Vint, Chemical Works, Seaham Harbour.

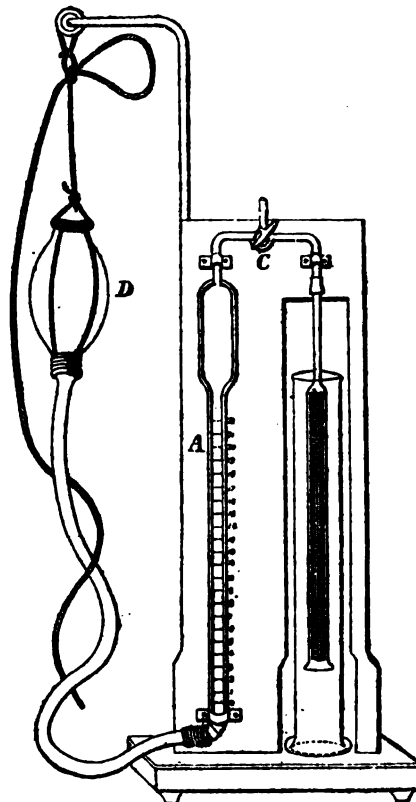
The CHAIRMAN, in drawing attention to the collection of apparatus sent for exhibition, alluded to the difficulty which had always been experienced in getting the meeting immediately preceding Christmas satisfactorily filled in the usual manner—a difficulty which had led to exhibitions of little novelties such as were then before the meeting, and to the passing the evening in a less formal and more social manner. The secretaries had been in correspondence with such friends as they thought likely to contribute objects of interest, including Messrs. Aug. Bel and Co., from whom they had been hoping till the last day to receive M. Cailletet's apparatus for the condensation of

gases, which in itself would have been ample attraction for the meeting; but, unfortunately for the Society, it was still in use at the Royal Institution, and in consequence could not be spared; but in the absence of this great attraction, there were yet objects that would repay examination, and he invited Mr. Starks to describe the mode of using the apparatus for estimating free oxygen in a mixture of gases, and the two forms of nitrometer.

The following notes, descriptive of the apparatus, were read:—

"*Apparatus for Estimating Free Oxygen Volumetrically,*" by Messrs. MAWSON and SWAN.

The accompanying sketch represents a simple and convenient apparatus for the rapid and correct volumetric estimation of oxygen when mixed with other gases. It consists, as will be seen, of a measuring tube, A, holding from 0 to the three-way tap 100 c.c., and graduated in the

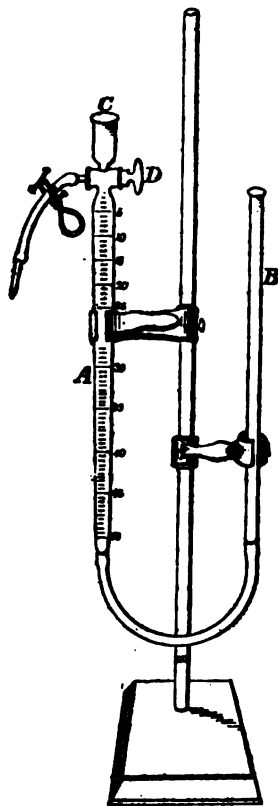


lower narrower portion into c.c. and tenths; of an absorption arrangement, B, which is a glass bell in which is placed a coil of copper gauze, suspended in an outer glass vessel which contains the absorbing liquid; a three-way tap, C, which serves to communicate between the measuring and absorption tubes, or with either and the external atmosphere; and of a globe, D, which serves the purpose of drawing the gas for examination into the measuring tube, and chasing it backwards and forwards between the measuring and absorption tubes. The solution used to absorb the oxygen is made of two parts of a saturated solution of chloride of ammonium and one part of liquid ammonia, s.g. 880. For an examination, the vessel, B, having been about two-thirds filled with the ammonium solution and sufficient water put into D to fill the measuring tube, expelling the air by the vertical outlet, C, the tap, C, is opened so as to make communication between A and B; D is then lowered so as to fill the glass bell and tube

above it is as far as the tap with the ammonium solution, then close c on the side b, and open the vertical outlet; d is then raised so as to fill the measuring tube with water up to the tap; the vertical outlet is now connected with the gaseous mixture to be examined, and by again lowering d the gas is drawn into the measuring tube to c, when exactly 100 c.c. will be present for examination; now turn c so as to make communication again between a and b, and chase the gas backwards and forwards by the raising and lowering of the glass vessel, d, two or three times, finishing by bringing the gas back to the measuring tube, and the absorbing solution to the original starting point; lastly, close the three-way tap altogether, and bring d down so that the two water levels in d and a coincide, when the reading of the liquid in a will give the percentage of oxygen that has been absorbed. By using other solutions it is obvious it may be used for the examination of other gases.

"Dr. Lunge's Nitrometer for Determination of Nitrous and Nitric Acids," by Messrs. MAWSON and SWAN.

The method of using is thus described by Dr. Lunge:—You will observe the measuring tube, a, holding 50 c.c., divided in $\frac{1}{10}$ ths, provided with a funnel, c, and a two-bore glass cock, b, hanging in an instantaneously-opening



spring clamp; an elastic tube connects a with the plain tube, b, of equal size, sliding up and down in another clamp. First of all, b is placed so that its lower end is nearly at a level with d; mercury is poured into b till it has filled a and has got into c; d is turned so as to close the communication between a and b, and so run off the excess of mercury through the lateral bore; b is lowered in its clamp, and 1 c.c. of the nitrous vitriol, &c., to be tested is introduced into c by means of an accurate pipette. By carefully turning d, the acid is sucked into a without any air following, and c is rinsed out by a few c.c. of pure acid, run into a in the same wa-

A is now taken out of the spring clamp, violently shaken for two minutes (no more is necessary), replaced in the clamp, and b adjusted, so that the levels of mercury of both tubes are equal, except that an allowance is made in b for the acid contained in a, $\frac{1}{10}$ th of the height of the latter being either allowed for as an addition to the mercury in b, or else being deducted from the barometrical pressure. After the temperature has become constant (quarter of an hour suffices for factory work, one hour for exact analyses), the volume of gas in a is read off (which can be done to $\frac{1}{10}$ c.c.) and reduced to 0° and 760 m.m. in the usual way. Each c.c. at 0° and 760 m.m. corresponds to 1.343 m.gr., NO = 1.701 m.gr., N_2O_3 = 2.417 m.gr., N_2O_5 = 3.805 m.gr., NO_2Na , &c. Now d is opened so that a and c communicate, b is raised, and thus first the gas then the acid are driven into c, from whence the acid is run away by the lateral bore of d; everything is now ready for a new test.

"Model of Modern Blast-Furnace," by P. A. BERKLEY.

MR. P. A. BERKLEY exhibited a model of a blast-furnace, which he said appeared to give the most satisfactory results in working, producing about three times as much iron per week as the old forms of furnace, and had not been surpassed by furnaces of either larger or smaller dimensions. It had worked incessantly since 1872. The model was accompanied by the following note:—

The model shows section through middle, with the various charges of iron ore, limestone, and coke, made to the scale of $\frac{1}{4}$ inch equal 1 foot. The counterpart of this furnace is at Jarrow. It is 85 feet high, 26 feet bosh—nearly 30,000 cubic feet capacity. It is blown by 5 tuyeres, 5 inches and $5\frac{1}{2}$ inches in diameter, with blast at a pressure of $4\frac{1}{2}$ lbs. and heated up to about 1200° F. The heat for the blast is raised from the waste gases from the furnace, as well as the steam for working the blast engine. This furnace is making 500 tons of pig-iron per week, in the making of which it consumes about 1750 tons of Cleveland iron-stone, or other ores equivalent thereto, 587 tons of coke, and 300 tons of limestone. The furnace was erected in 1872, for Messrs. Palmer and Company, by Mr. Philip A. Berkley, the Company's Engineer and Blast-Furnace Manager.

A brief discussion followed, in which Messrs. Hill, Morrison, Berkly and the Chairman took part.

(To be continued).

NOTICES OF BOOKS.

Ozone in Relation to Health and Disease. (An Address delivered before the Congress of the Sanitary Institute of Great Britain, held at Stafford, 1878.) By HENRY DAY, M.D. London: J. and A. Churchill.

IN this Address the author gives us the history of the discovery of ozone, and notices the successive theories of Schöenbein, Williamson, and Odling concerning its nature. He then describes the pathological action of this form of oxygen, and reveals facts which will probably startle those who believe ozone and "ozonised" articles of food or of medicine to be universally beneficial. He describes the death of animals after exposure to ozonised air under symptoms closely resembling those of acute bronchitis. He considers that if present in excess in the atmosphere, catarrh, bronchitis, and even pneumonia would be its natural results. Whether there is ever such an excess as would involve these consequences is an open question. He feels also bound to admit, according to the researches of Dr. Moffat, that during "ozone periods" apoplexy, epilepsy, vertigo, neuralgia, and diarrhoea are more frequent. Further investigations in this direction are imperatively needed, but what has been said may serve as a caution to dabblers in science who keep an ozone apparatus in action in their sitting-rooms as a prophylactic against diseases in general.

The absence or the deficiency of ozone has been, perhaps, too hastily placed in connection with zymotic disease. But that such a connection exists in case of cholera can scarcely be doubted. The author shows that in 1864 in the Bombay Presidency, cholera was in its greatest ascendancy when ozone was either wanting or at its minimum; that the disease showed a most marked decrease when ozone was registered as increasing, and when at its maximum the epidemic ceased altogether if the maximum continued for any time. Similar results were obtained at Strassburg in 1854 and 1855, and the experiments of Mr. Glaisher and of Dr. Moffat give confirmatory testimony. Whether there may be other causes in operation in addition to deficiency of oxygen is still doubtful. As a disinfectant the author pronounces it the best, safest, and least objectionable known. That it may kill disease-germs—whatever they may be—is no doubt highly probable from its action on the superior animals; but the question arises, Which will be killed first? We are somewhat surprised at finding in this address no reference to the well-known and justly admired work of Dr. C. B. Fox.

CORRESPONDENCE.

THE INSTITUTE OF CHEMISTRY.

To the Editor of the Chemical News.

SIR,—The long time that elapsed between the date of the Conference and the issue of the report to the members of the Institute, as alluded to by your correspondent "An Appeased Promoter," was caused chiefly by the decision to send it out together with the annual report of the Council.

The financial position of the Institute at the present time is excellent, but I fear few people would consider it to be such if the Council were to follow your correspondent's advice and "spend the large sum they have in hand," especially when it is remembered that our investment in consols does not cover the Life Compositions and Entrance Fees received up to the present. I do not for an instant suppose that "An Appeased Promoter" desires this to be spent. A man who lives up to his income is a poor man, however large his income may be, and this saying applies to corporations as well as to individuals. If, however, your correspondent thinks that the Council has left anything undone which it ought to have done, and will send me any suggestions, I can safely promise that they will have the most careful consideration from the Council.

In conclusion let me thank "An Appeased Promoter" for his frank reply, and at the same time to express a wish that he had not written anonymously, as it would have enabled me to have answered his first letter more satisfactorily, and have saved valuable space in these columns occupied by Yours, &c.,

CHAS. E. GROVES, Secretary.

Somerset House Terrace, W.C., March 22.

TENACITY OF STARCH.

To the Editor of the Chemical News.

SIR,—In reply to Mr. Thompson's letter (CHEMICAL NEWS, vol. xxxix., p. 122) I beg to say that the method for determining the tenacity of starch is not intended to determine the tenacity of a sample of British gum or the relative tenacities of British gum and starches, but only to compare samples of starches with each other. When I devised the method I conducted some experiments on the starches found in the market. I found that when equal weights of the different samples of starch were boiled with equal quantities of water and pieces of cloth of equal size starched with the different samples, that those samples which had the greatest tenacity gave the stiffest cloth.

Any of your readers can try this by procuring samples of Coleman's and Glenfield's starch weigh equal quantities, and boil with equal quantities of water and starch two pieces of cloth of equal size, *without squeezing out the excess of starch*, and dry; the touch will be able to determine which is the stiffest. Mr. Thompson says: "and weight for weight the farina which possesses the least 'tenacity' would produce the greatest stiffness, because the thinner the sample boils the more easily will the solid matter be able to enter the fibre, and it will enter in greater quantity." In the process of sizing when the yarn has passed through the size trough and got saturated with the size (say starch) it passes between two rollers, which squeeze out the excess of size. Now, although the thinner sample may enter the fibre more readily and in greater quantity than the more solid, yet in passing through the rollers the thinner sample will be squeezed out in greater quantity and more readily than the thicker sample, and as a consequence weight for weight the sample with the greatest tenacity will give the stiffest cloth, and what the manufacturer has found out by experience is correct in theory, and Mr. Thompson and not my reasoning is fallacious.—I am, &c.,

GEORGE WHEWELL.

Exchange Chambers, Blackburn,
March 25, 1879.

TESTING OF GAS LIQUORS.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxviii., p. 193, there is a short article by Mr. T. H. Davis, "On the Testing of Gas Liquors." To those who have many of these liquors to test any contrivance that would tend to abridge the labour and at the same time afford reliable results would be hailed as a boon, and Mr. Davis is much to be commended for his efforts in that direction. The usual commercial valuation of gas liquors by degrees in Twaddle's hydrometer is so unsatisfactory that it is quite time it was superseded by some better method, and I think the test proposed by Mr. Davis gives at any rate a more reliable indication of the real strength of gas liquor than the method usually employed. I have not, however, found that this test is sufficiently accurate to supersede the usual distillation process, and I am not sure that Mr. Davis intends it should do so.

I have recently had five gas liquors placed in my hands for examination, and below I give the results of analysis by the method proposed by Mr. Davis (A), and also by the process of distillation (B)—

NH₃ per 100 parts of the Liquor.

		A.	B.
No. 1		2.55	2.93
" 2		2.24	2.66
" 3		1.59	1.86
" 4		2.12	2.72
" 5		2.38	2.84

I am, &c.,

A. McDONALD GRAHAM.

Wellfield Villa, Turnchapel, Plymouth,
February 22, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 9, March 3, 1879.

Reply to M. Van Tieghem Concerning the Origin of the Amylobaeter.—A. Trécul.—The author points out that his words have been inaccurately quoted by M. Van Tieghem.

Emissive Power of Coloured Flames.—M. Gouy.—A mathematical paper, not susceptible of useful abstraction.

Absorption Spectra of Didymium and of Certain other Substances Extracted from Samarskite.—J. L. Soret.—The author has compared didymium chloride extracted by M. Marignac from samarskite with a sample from a different source. In the entire less refrangible part of the spectrum, from the red to the green, he finds no appreciable difference either in the colour or in the intensity of the rays, and the spectrum agrees very exactly with the figures in the work of M. Lecoq de Boisbaudran. In the blue and the indigo differences of intensity appear. Hence the author is led to the conclusion that the didymium from samarskite contains some foreign body. The same substance is present to a smaller extent in terbia, and less still in the didymium obtained from cerite.

Action of Ammonium Sulphocyanide upon Monochloric Aceton.—T. H. Norton and J. Tcherniak.—Instead of sulpho-cyan-aceton, as was expected, there was produced the sulphocyanide of a base, $C_4H_5N_2S$.

Amidated Acids Derived from the α -Butyric and Isovaleric Acids.—E. Du villier.—The acids described are the methyl-amido-isovaleric, the ethyl-amido- α -butyric, the ethyl-amido-isovaleric, the phenyl-amido- α -butyric, and the phenyl-amido-isovaleric.

Influence of Oxygen on Alcoholic Fermentation as Produced by Beer Yeast.—A. Béchamp.—Oxygen was found to exercise a favourable influence upon the production of alcohol. The quantity of acetic acid seems to depend much more on the temperature and on the nature of the yeast than on the oxygen. In short, the oxygen seems to act as a stimulus, under the influence of which the life of the yeast and the changes of its matter are more active. In a second set of experiments it was proved that the oxygen was actually absorbed.

Chemiker Zeitung.
No. 2, 1879.

Detection of Chromates and Free Chromic Acid.—To detect mono-chromate along with bichromate add to a few c.c. of the boiling solution a drop of a moderately concentrated solution of manganese sulphate. In presence of monochromate a blackish brown precipitate is formed. Bichromate in monochromate is detected by adding to a boiling solution of hyposulphite of soda an equal volume of a hot solution of the chromate in question. A brown precipitate or a distinct turbidity proves the presence of bichromate. The precipitate is chromic peroxide. Free chromic acid in a solution of bichromate is detected by adding solution of iodide of potassium and agitating with sulphide of carbon, which is coloured a deep purple by the liberated iodine.—*Zeitschrift Anal. Chem.*, 18, 78.

Determination of Sulphites and Hyposulphites.—If both salts are present together in solution, it is necessary to find the quantity of iodine which a part of the liquid requires if mixed with acetic acid; and the quantity of sulphate of baryta which an equal part of the solution yields after complete oxidation with bromine. Two equations are thus obtained with two unknown quantities. If a sulphate is also present its quantity is ascertained by adding to the liquid bicarbonate of soda, passing a current of carbonic acid through the liquid, heating after expulsion of the air, and after the addition of hydrochloric acid in excess concentrating to quarter its volume. The sulphurous acid being thus expelled the sulphur is filtered off, and the sulphuric acid in the filtrate is determined in the ordinary manner.—*Zeitschrift Anal. Chem.*, 18, 79.

Separation of Gold and Silver.—The metal is alloyed with 5 to 8 parts of zinc, for which the heat of a Bunsen burner is sufficient. The alloy is dissolved in nitric acid, when gold (with platinum and tin as stannic oxide if present) remains undissolved. To separate gold from platinum and tin they are dissolved in aqua regia, the

platinum metals are precipitated with ammonia, the free chlorine is expelled, and the liquid is mixed with excess of ammonio-ferrous sulphate, the excess of which is determined by titrating back with permanganate.—*Zeitschrift Anal. Chem.*, 18, 104.

Reactions of Bile Acids and their Detection in Urine.—A solution of the bile acids, if oxidised by the addition of ferric chloride, antimonie chloride, lead peroxide, barium peroxide with the addition of hydrochloric or sulphuric acid, especially if exposed to direct sunlight, passes through a variety of colours from yellow, red, vinous red, to blue and blue-violet. Similar colours are produced by the addition of stannous or antimonious chlorides along with sulphuric acid. Casali takes advantage of this play of colour for the detection of bile acids in urine. The urine is precipitated with sugar of lead and ammonia, the precipitate treated with ether and dilute hydrochloric acid, and the ethereal extract is drawn off and evaporated in three porcelain capsules at common temperatures. To the first residue are added barium peroxide and sulphuric acid; to the second tin crystals and sulphuric acid; and to the third antimonious chloride and sulphuric acid.—*Zeitschrift Anal. Chem.*, 18, 128.

Examination of Water for Bacteria.—Himly fills a flask, previously washed out first with hot concentrated sulphuric acid and then with the water in question, three-quarters full of the water, and adds a little solution of meat-extract, prepared by boiling extract for fifteen minutes in a large quantity of water. The flask is then stoppered and heated to $98^\circ F.$ on the sand-bath. Bacteria, if present, increase rapidly, occasioning a turbidity and a film on the surface of the water. The experiment requires about twenty-four hours. Distilled water similarly treated shows no turbidity.—*Zeitschrift Anal. Chem.*, 18, 117.

Action of Animal Charcoal on Salts.—Many salts, on filtration through animal charcoal, are held back more or less completely, or decomposed so that a part of the acid runs through, while the rest, in combination with the whole of the base, is retained by the charcoal.—*Zeitschrift Anal. Chem.*, 18, 97.

No. 3, 1879.

Dr. J. Volhard has been appointed Professor of Chemistry at the University of Erlangen, as successor to the late Gorup-Besanez.

A. Podewils proposes to convert faecal matters into poudrette by treatment with smoke, either of wood or coal.

Determination of Ferrous Oxide in Silicates.—Döller recommends the following process:—The finely pulverised mineral is covered with sulphuric acid and hydrofluoric acid in a platinum crucible set upon an iron plate. Over the plate is inverted a high beaker, whose edge fits into a groove made in the margin of the plate, and made air-tight by means of mercury or sand. From above, through a hole in the bottom of the beaker, a tube is passed to just over the crucible, and introduces a constant stream of carbonic acid. The whole apparatus is heated on the sand-bath, and the hydrofluoric acid escapes through the hole in the bottom of the beaker.—*Zeitschrift Anal. Chem.*, 18, 50.

No. 4, January 23, 1879.

A Contribution to the Fat-Industry.—Dr. B. Terne.—The author speaks of the application of the vapour of benzol in America for extracting oils and fats, not merely from wool and offal, but from bones, hoofs, &c. The fat of swine he remarks is apt to contract an objectionable colour. The chief difficulty is the removal of the characteristic odour of benzol. He has tried the action of benzol vapour upon tallow-greaves which had been previously submitted to hydraulic pressure of 10,000 to 12,000 lbs. per square inch, and obtained from 8 to 10 per cent of fat. The new Sanitary Office at Gera carefully examines all meat for *Trichina*.

Dr. Upmann, in consequence of some experiments upon

dogs, pronounces oxalic acid harmless. Dr. Pfeiffer, in reply, points out that the stomach and intestines of dogs generally contain considerable quantities of calcium phosphate by which the oxalic acid would be converted into calcium oxalate, an inoffensive salt.

According to the *Revue Industrielle* a metal has been extensively produced in France, which is a combination of iron and steel. The two metals are run separately into a mould divided into two parts by a thin plate of sheet iron, when the whole becomes welded together. The new metal is particularly adapted for armour-plating, anchors, and safes.

According to Fischer, in the *Metallarbeiter*, petroleum is adulterated with so-called solar oil, which occasions the dull yellow colour of the ordinary oil and its yellowish flame.

No. 5, January 30, 1879.

Dr. H. Geissler, of Bonn, died on the 26th ult. His establishment for the construction of physical apparatus will probably be carried on by Franz Müller, who has been for twenty-five years first his pupil and afterwards his assistant.

Lead, in dangerous quantities as an ingredient in the so-called tinning of sauce-pans, &c., has been detected by Dr. Brockhoff, of Magdeburg.

Dr. Wolff, inspector of factories for the Düsseldorf circle, has introduced a respirator for the use of men employed in white-lead works, lead smelting establishments, &c. The results are said to be satisfactory.

Test for Olive Oil.—Poutet proposes the following method:—Nitrate of mercury (doubtless mercurous) is prepared by dissolving 6 grms. mercury in 7.5 grms. nitric acid at 38° to 46° B. in the cold. Next 96 grms. of the oil in question are mixed with 8 grms. of the mercurous nitrate and shaken together every ten minutes for two hours. After being allowed to settle for twelve hours, if the oil was pure the elaidin thus formed is pale yellow and quite solid. In adulterated samples the elaidin is orange or dark red, and only partially or not at all solid. To detect the presence of the oil of sesame 2 parts of the sample are shaken up at a temperature of 20° to 25° with 1 part of pure hydrochloric acid at 22° B., in which 0.05 to 0.1 grm. of sugar has been previously dissolved. After standing for some time the oil separates from the acid, and if sesame is present assumes a rose colour. The more intense this colour the greater is the quantity of the impurity.

No. 6, February 6, 1879.

A. G. Moser, of Graben, in Thun, is said to have invented a remedy for the phylloxera, which completely eradicates it without injury to the vines.

T. Salzer has observed that sal-ammoniac and calcic hypochlorite react upon each other so violently as to give rise to an explosion.

Schneider, on preparing sodic hypochlorite by passing chlorine-gas into sodic bicarbonate, observed a rose colouration, which he traces to the presence of sodic ferrate.

No. 7, February 13, 1879.

A curious toxicological case is reported from Hamburg. The body of a man who died in 1867 was taken for examination. It was thought necessary to determine arsenic not merely in the corpse in question, but in the soil of the churchyard at different distances from the coffin, and also in the body of another man who had been subsequently buried in the same grave. This latter body was perfectly free from arsenic, which, however, was found in the first corpse in ample fatal quantity (0.24 grm.), whilst in the lid of the coffin and in the adjacent very minute quantities were traced. Hence the conclusion was fairly drawn that the man in question had been poisoned with arsenic, and that a portion of the poison had been gradu-

ally transferred from his body to the wood of the coffin and the adjacent soil.

Determination of Iron in Grain and other Food-Plants.—Eliosoff proposes the following method for the determination of iron in presence of phosphoric acid. The ash is evaporated to dryness with nitric acid, again taken up in dilute nitric acid, filtered, concentrated, precipitated with the molybdate of ammonia; the precipitate is digested upon the filter for some time with citric acid, the filtrate evaporated to dryness, the residue ignited, and re-dissolved in hydrochloric acid with the aid of heat, and the liquid separated by filtration from the liberated molybdic acid. In the filtrate the iron is determined as usual.—*Journ. de Pharm. et de Chimie*.

No. 8, February 20, 1879.

Dichromates.—Schulerud has prepared the dichromates of silver, thallium, and lithium, but has not succeeded in obtaining corresponding salts of lead, barium, and mercury. He concludes that univalent metals only yield dichromates, and argues hence to the univalence of lithium.

Detection of Salicylic Acid in Beer.—Blas recommends to drink the beer and in about three hours to test 20 c.c. of the urine with ferric chloride in the usual manner, as the reaction is about five times more delicate in urine than the original beer.

Extraction of Animal Pigments.—C. Méhu separates colouring matters from animal fluids by the addition of ammonium sulphate. In this manner he readily isolates the pigments of pathological urines, fecal matter, bile, serous liquids, &c.

Formation of Mannite in Beer.—The formation of mannite is always a mark of the want of cleanliness, and is particularly promoted by decaying wood. In such cases the sugar present enters not into the alcoholic but the mucic fermentation.

Moniteur Scientifique, Queneville. January, 1879.

Critical Examination of a Posthumous Work by Claude Bernard on Alcoholic Fermentation.—M. Pasteur.—This is merely the reproduction in full of a paper read before the Academy of Sciences, Nov. 25, 1878.

Causes of Death in Carbuncular and Septicæmic Affections.—M. Colin.—According to the author's views there is a death by the blood resulting from the inaptitude of this liquid to maintain the life of the cells and of the other anatomical elements. This type of death must be added to those characterised by Bichat, and is probably common to a great number of maladies, the carbuncular, the putrid, the typhic, and the pestilential.

Chemical Composition of Yeast.—M. Nægeli.—In this lengthy paper the author ascribes to bottom-yeast the following chemical composition:—

Cellulose and vegetable mucilage, forming the cellular membrane	37
Proteic matters—(a) In the state of albumen ..	36
(b) In state of phosphorous compounds ..	9
Peptones, precipitable by sugar of lead ..	2
Fatty matters	5
Ash	7
Extractive matters, &c.	4

100

Under the extractive matters are included small quantities of invertine, leucine, and grape-sugar; still smaller quantities of glycerin, succinic acid, cholesterin, guanin, xanthin, sarkin, probably of inosite and traces of alcohol.

Alkaloids of Veratrum.—A. Tobien.—Taken from the *American Journal of Pharmacy*, 1878, No. 3, p. 12

Optical Rotatory Power of Liquids and Dissolved Substances.—O. Hesse.—Translated from the *Annalen der Chemie*.

Action of High Temperatures upon Petroleum, Coal-tar, and Analogous Substances.—Alexandre Letny.—The author finds that the more elevated the temperature of the furnace which heats the still, the more the distillate is resinous and difficult to purify. This resinification becomes still more sensible when the flues are higher than the level of the petroleum in the still, in which case the vapours are decomposed by their contact with the heated sides. In the distillation of coal-tar and ozokerite, where it is necessary to avoid decomposition, the still-head and the pipe for the escape of vapours are placed as near as possible to the liquid; otherwise the paraffin would be decomposed. At augmented pressures and at a red heat petroleum is resolved into gaseous compounds.

On Ultramarine.—Dr. R. Hoffmann.—Taken from *Liebig's Annalen*.

Poisoning by Mushrooms.—J. A. Palmer.—Mushrooms may act as a poison in three different manners. They may act as an indigestible matter, which is the case with hard, coriaceous species, and may even occur with the edible mushroom when decomposing, as it gives off sulphuretted hydrogen in quantity sufficient to cause vomiting. Or, again, they may be glutinous or acrid. Many Boleti, otherwise innocent, are too gluey to serve as food. Lastly, mushrooms may contain a subtle alkaloid devoid of smell and taste, as happens in the group of the Amanitæ. This compound is known by the name of Amanitin, and to it the fatal cases of mushroom-poisoning are mostly due. No remedy has yet been found. No immediate effects are produced by this poison, but after eight to fifteen hours the patient experiences stupefaction, nausea, and diarrhoea, followed by delirium and death. Mushrooms containing this poison seem able to communicate it to wholesome species by contact, and it may also be absorbed through the skin. The author was on one occasion seized with alarming symptoms after carrying in this hand some Amanites wrapped in paper.

Resists in Calico-printing.—A. Schultz.—A purely technical paper, incapable of useful abstraction.

Aniline-black upon Wool and other Textiles.—M. Delory.—The author first mordants in 100 grms. each of potassic bichromate and oil of vitriol dissolved in 10 litres of water. The dye-bath he prepares by dissolving 30 grms. aniline hydrochlorate in 9 litres of water at 28°. In another vessel he dissolves 55 grms. potassic bichromate in 1 litre of boiling water, and adds to the hot solution 48 grms. sulphuric acid at 66° B. The solutions are then mixed together. In this liquid the tissues are worked for an hour without heat; the temperature is then raised quickly to 95° to 100°, and kept at that heat for 25 to 30 minutes. At the moment of applying the heat 10 to 12 grms. sulphate of copper are added, previously dissolved. The material is then taken out, washed very well, and taken through an alkaline bath of soap and soda, to which have been added 0.2 to 0.5 grm. aniline-violet per litre of water.

Historic Notes on the Discovery of Dyeing with Aniline-black.—In these notes M. Girardin, whilst admitting Mr. John Lightfoot's claim as the original discoverer of aniline-black, maintains that a M. Carl Stalare, of Lille, was the first who successfully used the new colour in dyeing under the name of "French black."

Poisonous Character of Salts of Copper and the Use of Copper Sulphate in Panification.—A collection of opinions on the innocence of the salts of copper.

February, 1879.

Industrial Society of Mulhouse.—The hydrosul-

phurous acid of Schützenberger and Lalande has been proposed for bleaching animal fibres in place of sulphurous acid. The white obtained is, however, not pure.

The insoluble ferricyanides are capable of oxidising indigo without the aid of an alkali.

Prof. Kopp communicates spectroscopic observations on eleven derivatives of resorcin, which he has just examined. The plates which accompany his memoir, and which represent the spectra of these different compounds, indicate the rays which he considers characteristic of methyl, bromine, iodine, &c., and which are the basis of his analytical method. M. Kopp, having as yet no data on the composition of these products, desires to obtain from the manufacturers who have furnished them the verification of his analyses before the publication of his memoir.

Colouring Matters Belonging to the Group of Azoic Bodies.—A. Kopp.—We have here an account of the preparation and properties of Bismarck-brown, of Magdala red, of azo-diphenyl blue, safranin, the chrysoidins, tropæolins, oranges, &c.

Glasses Melted with Alkali Alone.—Dr. P. Ebell.—Sodic and potassic sulphides, or sulphates in presence of reducing agents, on dissolving in glass at the melting-point impart to it a peculiar tint, varying from yellow to brown, and even to a deep red-brown if in sufficient quantity. In certain conditions the same colouration is produced by the introduction of free sulphur into glass, melted and kept at a high temperature. This colouration takes place only in glasses where there is sufficient available alkali to give rise to sulphides. To produce the colour there must be at least 1 equivalent of base to 2.5 of silica. The base may be lime or baryta, as well as potash or soda. Silicates in igneous flux are solvents for very various bodies, simple or compound. They dissolve free metals, such as gold, copper, silver, and lead; oxides, like chrome, alumina, stannic oxide, magnetic oxide of iron; also salts, such as sulphates, phosphates, aluminofluorides. These bodies are separated out on cooling either in a crystalline or an amorphous state, according to circumstances, and communicate different properties to the glass. The affinity of silica for bases measured by the carbonic acid displaced during fusion is not a constant magnitude, but depends on an action of mass. The quantity of carbonic acid expelled by a given weight of silica is so much the smaller the less alkaline carbonate there is in reaction.

Aniline and the Methylated Toluydins and the Colouring Matters thence Derived.—P. Monnet, F. Reverdin, and E. Noelling.—Already noticed.

MEETINGS FOR THE WEEK.

MONDAY, 31st.—Medical, 8.30.

TUESDAY, 1st.—Royal Institution, 3. "Animal Development," Prof. Schäfer.

— Zoological, 8.30.

— Society of Arts, 8. "Remarks upon an old Map of Africa Contained in Janson's Atlas, Published in Paris in 1612," and exhibited by R. Ward. "The Submarine Telegraph to South Africa," by J. Sivewright. (African Section.)

WEDNESDAY, 2nd.—Society of Arts, 8. "Some Causes of the Recent Depression in Trade," by B. Francis Cobb.

— Pharmaceutical, 8.

THURSDAY, 3rd.—Royal, 8.30.

— Royal Institution, 3. "Sound," Prof. Tyndall.

— Royal Society Club, 6.30.

— Chemical, 8. "On Terpin and Terpinol," Dr. Tilden. "On the Transformation of Aurin into Trimethyl-para-rosanilin," R. S. Dale and C. Schorlemmer. "On a Gold Nugget from South America," A. H. Wood. "On the Solution of Aluminium Hydrate by Ammonia and a Physical Isomeride of Alumina," G. C. F. Cross.

FRIDAY, 4th.—Royal Institution, 9. "Molecular Physics," by Mr. Crookes.

— Geologists' Association, 8.

SATURDAY, 5th.—Royal Institution, 3. "Etching," by Mr. Seymour Haden.

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BLUE FLAME FROM COMMON SALT.

By A. PERCY SMITH, F.C.S., F.I.C.

SOME time ago the question was raised in *Nature* concerning the origin of the blue flame produced when common salt is thrown into a hot fire.

Among the suggestions that were advanced, no one offered the only explanation that is at all feasible, viz., that it is due simply to hydrochloric acid.

The blue flame is not produced by sodium chloride only, but by other chlorides as well. Those I have tried are— BaCl_2 , SrCl_2 , KCl , AmCl , Hg_2Cl_2 , and HCl , the last both in solution and as gas. It would be waste of time and space to enumerate all the experiments I have made on this subject; many of them were for the purpose of proving that neither carbon nor sulphur had any share in the reaction.

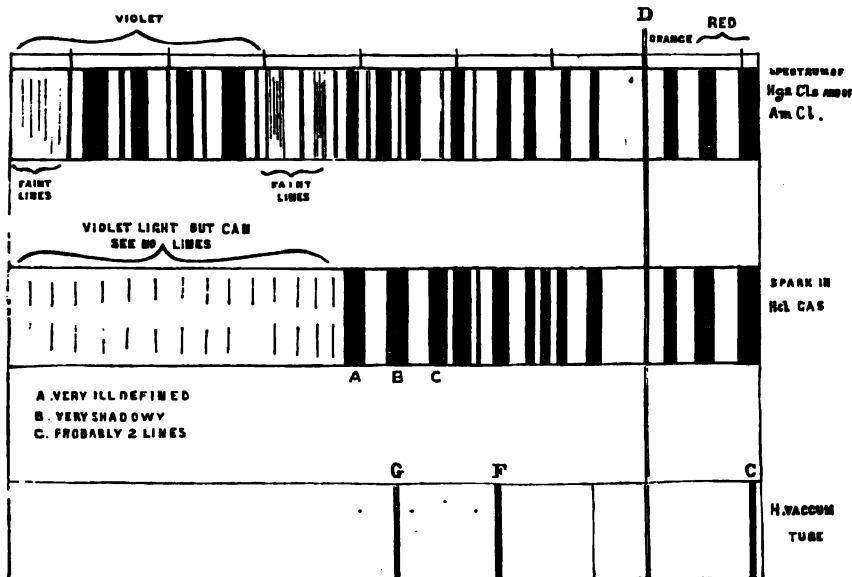
do not pretend to any high degree of accuracy in my delineations of the lines; besides the different conditions under which the two were compared might account for considerable variations.

For the HCl I used a coil (capable of giving a 2" spark) with a Leyden jar, and worked by 6 Smées. The carbon points were $\frac{1}{8}$ -inch apart. The spark was tried both focussed and unfocussed on the slit of a 2-prism spectro-scope.

When comparing the two spectra side by side it was difficult in some cases to be sure of coincidence, because the flame from the Hg_2Cl_2 would flash out brilliantly for a moment and quite overpower the more feeble lines of the HCl ; it would then disappear entirely, and more chloride would have to be placed on the gauze.

I have no doubt in my own mind that HCl is the cause of the blue flame. I have proved that Cl is a necessary constituent, and I have not been able to get it in the absence of hydrogen (the spectrum of pure Cl is very different); and, besides, the red H line is present in both cases, and probably the other two as well. I do not think that the presence of aqueous vapour is sufficient to account for the red line.

I subjoin the spectra as I mapped them; but it must be borne in mind that I do not vouch that they are without error. I intend to photograph them when I have sufficient



One of my methods of obtaining the flame was to burn pure hydrogen from a glass jet, and allow a mingled stream of HCl and NH_3 from two other jets to pass into it. The best source, however, is calomel, heated on wire gauze by a Bunsen burner; the next best AmCl . The spectrum of the "chloride" flame is characterised by a series of double bands in the green, blue, and violet, the least re-frangible of each pair being the broadest. The four pairs in the violet are especially prominent. There are two red bands or lines and one orange. The least re-frangible red line occupies the place of the hydrogen line C. A spark between carbon points in a bottle of HCl gas yields a spectrum similar in appearance to that obtained from a chloride, but I was unable to see any violet bands, only a faint continuous spectrum. I was able to ascertain that the red lines coincided exactly, but I cannot affirm with the same positiveness that all the green lines and bands coincide. Some undoubtedly do; but the spectrum of the HCl was so faint and the spectrum of the chloride so transient that measurements were very difficult, and I

leisure, and I hope the results will be more definite. I may be able to find violet lines in the spectrum of HCl .

Rugby, March 15, 1879.

ON SUPERSATURATED SOLUTIONS.

By J. G. GRENFELL, F.G.S.

I STAND corrected by Prof. Tomlinson thus far, that his paper was *received* by the Royal Society in September, 1877, and not *read*, as I previously stated.

As regards my description of his paper as one intended to upset my theories and uphold his own, I cannot see that Prof. Tomlinson has to find fault with. In using those words it never occurred to me that any one would emphasise the pronouns *my* and *his*, and suppose that I intended to imply he had any personal feeling in the matter. No one who knows Prof. Tomlinson would sup-

pose that for a moment. He battles stoutly against what he believes to be error, and I found no fault with that. What I did complain of was the passage which I quoted in my letter of January 10th. Every one to whom I have shown his paper agrees that the words as they stand distinctly imply that I denied the fact of the lecture-table experiment with Glauber's salt.

Prof. Tomlinson denies that he anywhere implied this. I am glad to accept this as an expression of his intention, but I cannot admit it as a fair description of the words he used. I have laid stress on this only because the line taken by Prof. Tomlinson in that paper is that I am at variance with all my predecessors, a point to which I shall have to recur, and that after repeating some of my experiments his results were contradictory of mine.

Prof. Tomlinson also protests against my making him out to be ignorant of the very simple conditions necessary for the success of the experiments with drops on a glass plate. The case stands thus:—In his paper he gives a general statement, applicable apparently to all supersaturated solutions, that when a point was drawn through a drop so as to deform it, crystallisation sets in. This is in dry weather. I have drawn a point through hundreds of drops of the strongest solutions of sodium sulphate, carbonate, and acetate, on recently washed plates, in all kinds of weather, without causing crystallisation; I have just tried this with sixty drops of a strong solution of the acetate, and it was inactive in every case. I have deformed drops of ammonia alum, which had begun to deposit the modified salt, on a plate over calcium chloride, and repeated this with sodium carbonate and acetate over sulphuric acid, and the scratching had no effect whatever when the plates were put back over the chloride and acid.

I explain this difference in our results by supposing that Prof. Tomlinson did not take sufficient care in cleaning his plates. There is nothing which need hurt him in the supposition, because it is only the experience given by a long series of experiments with plates which has taught me what precautions have to be taken. It is in the same spirit that Prof. Tomlinson quietly dismisses what he calls my minute criticism, as throwing no light on the circumstances under which these solutions crystallise. Experience has taught me that it is only by the minutest possible criticism of each case that we can hope to arrive at the explanation of these very difficult phenomena. I recognise at present three well-established ways in which a supersaturated solution may be made to give the normal salt. The first is by introducing a crystal of the same salt, or of an isomorphous one of similar constitution as Mr. Thomson has recently shown. The second is by the mechanical action of absorbent substances, when not at once saturated by the solution. The third is by cold. Until it is proved that these three do not apply to the case in point, I refuse to look for any other cause.

Now the first two of these causes exhibit the most extraordinary delicacy in their mode of operation. Mr. Liversidge agrees with me in explaining the action of the minute aerial nuclei by their absorptive power. Hence if Prof. Tomlinson's plates had minute organic particles on them in dry weather these would cause the crystallisation of his drops when, by deforming them, he brought the solution in contact with these nuclei. He notices himself that if the tail formed sprang back crystallisation did not ensue, and I have always found that plates exposed to air are much more likely to be active than plates recently washed or dried under cover.

The extraordinary delicacy of the second cause has been proved in the course of my own experiments, of which I will give a few. A large glass plate, on which drops of sodium acetate had been made to crystallise, was carefully washed in cold water, and then dried and exposed to the air. A great many drops of a strong solution of the acetate were then put on it, and all remained liquid; but I found that scratching with a pin inside the drops made every one of them crystallise instantly, although the

every experiment. I naturally thought at first it was a case of scratching causing crystallisation; but I had so often scratched the acetate without effect that I was very loth to believe it. I then washed the plate with boiling water, and then scratching had no effect whatever on many drops of the same solution. The acetate has a very remarkable power of adhesion, and was still present in the depressions of the plate covered by weaker solution, but was unable to determine the crystallisation of the drops until scratching brought it in contact with the strong solution.

A few days ago I took an apparently clean plate, washed it carefully in cold water, put drops of the acetate on it, and then put it over calcium chloride. In an hour or two the modified salt was forming on the drops by evaporation; then scratching with a pin made every drop crystallise. This looked very like a confirmation of Prof. Tomlinson's results. But on washing the plate in hot water, and repeating the experiment, scratching had no effect whatever, and when put back over sulphuric acid for two days only the modified salt was formed. Probably the plate had previously been used for the acetate; possibly, in the first case, nuclei had not been thoroughly removed, but the acetate is so little sensible of absorbents that this latter supposition is not at all probable.

Again, a pin had been frequently used for making the acetate crystallise, and often heated red hot; it was rubbed through the fingers and then left $3\frac{1}{2}$ hours in cold water; it was then immediately active, not on quietly putting it into a drop, but on scratching; whilst boiling water cleaned it at once.

It is evident that these facts must be taken into consideration if we do not wish to have results ascribed to all sorts of wrong causes. Another fact of the same kind is not without interest. Dr. De Coppet has described two isomeric modifications of anhydrous sodium sulphate (*Comptes Rendus*, lxxviii., p. 194), one formed by dehydration above 33, the other formed at lower temperatures. This latter kind, he says, determines the crystallisation of a solution of the salt as the normal, whilst the former does not. M. Gernez suggested that this was because at a low temperature the sulphate did not perfectly effloresce. I tested the point by rubbing some of the salt very fine in an agate mortar and then exposing it for weeks over calcium chloride. I then took a plate with a large number of drops of the sulphate and tested various portions by taking up a little with a pin and introducing it into a drop. I found that all the thicker parts were immediately active, but that here and there a small very thin patch was inactive, and the anhydrous could be seen to dissolve in the drop. Hence it is clear that M. Gernez's explanation is correct. This method of testing is one of extreme delicacy for the presence of salts which form supersaturated solutions.

Another illustration of the necessity of minute investigation is the following:—

Some years ago I showed that scratching a solution of sodium sulphate in about six parts of sulphuric acid causes crystallisation. I attributed it at the time to vibration upsetting the state of unstable equilibrium. My colleague, Dr. Tilden, found that the same thing happened when drops of potash alum were scratched; and at a lecture in Bristol he gave it as another instance of vibration. But I have since found that by taking extreme care in cleaning the plate and using one recently washed scratching is by no means always active. I attribute this case also to the presence of absorbents on the plate. I am also strongly inclined to believe that the effect of scratching the acid solution mentioned above is due to the same cause. Rubbing drops of this with a hard, compact wood has no effect, but it always crystallises by absorption on the end of a match simply dipped in the drop and laid aside. Little shavings of a match kept floating on a drop act in the same way, but very often not if they are pushed into the drops. It is easy to see how the solution would gradually act on the organic particles

in the depressions of the plate, and then scratching brings the crystals which are forming into contact with the stronger solution. I did once find a plate inactive when scratched, but I have never been able to reproduce the conditions, although I have tried several ways of cleaning the plates. It must also be carefully borne in mind with regard to these experiments with drops that the sensitiveness to absorbents depends on the strength of the solution, and also on the temperature.

In hot weather I found that a rather weak supersaturated solution of sodium sulphate was absorbed by filter-paper without giving the normal salt even when dry, as the filter-paper was cut up and introduced into drops without producing any effect. This can never be done, as far as my own experiments go, with strong solutions of that salt. Absorbents also, such as filter-paper, have no effect whatever on sodium acetate except in the very strongest solutions.

It appears, then, that in discussing the question why these solutions crystallise or do not crystallise in a given case, we have to take account of the strength of the solution, of the temperature, of the possible presence of the salt when least expected, of the freedom of the plate from dry absorbent bodies derived from the air or elsewhere, and of a specific sensitiveness to absorbents peculiar to such kind of salt. This necessitates minute criticism, and with all deference to Prof. Tomlinson's opinion, I maintain that one or other of these considerations has frequently led me to a simple explanation of results which are otherwise most puzzling. The action of oil on these solutions is a very simple one according to Prof. Tomlinson; when a film adheres to the solution the normal salt separates. But I find that the phenomena are much more complicated.

When solution of sodium carbonate is dropped into oil, or is rubbed with oil, a salt separates at once, which is not the normal salt, as it can be introduced into drops without causing crystallisation. I have also seen a very pretty thin flexible opalescent film form at once round the sides of a drop of ammonia alum on oil. As I write, three drops of a strong solution of potash alum are lying unaltered in castor oil; yet I have often found oil rubbed on a plate powerfully unclear to this solution. On sodium acetate I have never found oil have any effect, nor on the sulphate when proper precautions are taken. It is quite possible that the oil acts by catching up absorbent particles of dust, but the subject needs investigation.

I have omitted to notice a property possessed by these solutions which may easily prove a source of error. When small *well-shaped* crystals of the salt are forming in a solution or in a drop they may continue to grow for hours very slowly; but if one of these is broken the drop crystallises with a flash. It may, as far as my experiments go, be taken as a general law that broken crystals have much more effect in producing crystallisation than perfectly-shaped ones. The crystal seems to have the power of growing with much greater rapidity along one of its axes than along the others. It is this property which makes a strong solution crystallise in fibres, while a weak solution gives well-shaped crystals. In the former case the growth of the more powerful axis does not sufficiently diminish the strength of the solution in front of it to check the rate of growth; in the latter case the crystal is surrounded constantly by impoverished solution, which gives time to the lateral forces to develop themselves.

Hence a drop on a plate may really be crystallising in the depressions of the plate, but very slowly, and the experimenter will find that scratching causes crystallisation at once, or that the drop will crystallise spontaneously after a certain time.

I find that a pin which has been heated red-hot often, and is used for making the acetate crystallise, and is then left in water for an hour or two, makes drops of the acetate crystallise very slowly in large well-formed crystals, whilst scratching makes it crystallise at once in fibres all over. This subject of the relation of crystalline form to

the conditions under which the crystals were formed is interesting to geologists as well as chemists.

The last point I have to notice is one which Prof. Tomlinson insists upon in his paper, and in his note to you; it is that I am at variance with all my predecessors. He says that Loewel, Violette, and Liversidge expressly state that porous bodies, bodies greedy of water and capable of hydration, are incapable of determining the solidification of these solutions.

He has forgotten, as I did at the time of making my experiments on absorption, that Mr. Liversidge, in his last paper in August, 1870, enunciated the absorption theory in these words:—

"(1) It is not impossible that nuclei consist of microscopic organisms which act by abstracting water, the action being set up at a point from which it is propagated throughout the mass; (2) that they are rendered inactive by heat because it entirely destroys them; (3) that their action is arrested by previous saturation in water, because they then can no longer abstract water from a saline solution."

He had only made a couple of experiments with lycopodium, which he found to be active when dry and inactive when wet, but he expressed a hope to have shortly the results of other experiments. Unfortunately, he never resumed the subject. My own experiments on absorption were begun five years later, when I had forgotten all about Mr. Liversidge's suggestion, and when I was working at a different part of the subject. They entirely confirm Mr. Liversidge's views as to the action of nuclei; they show that the action of absorbents in considerable quantity resembles that of nuclei, but with two important modifications, which determine the conditions under which the absorbents act—namely, that they are inactive if saturated at once by the solution, and that very rapid absorption often prevents crystallisation. It is only in virtue of this extension and modification that I can lay any claim to the theory as my own, and my results are in perfect harmony with those of the only person who, with the exception of Prof. Tomlinson and M. Gernez, has really worked at this subject in recent years. It is true that Loewel, Violette, and Liversidge state that phosphoric anhydride and other dehydrating substances have no effect; but I have shown that when these substances are introduced into a considerable bulk of the solution, so as to be saturated at once, crystallisation does not ensue.

ON INDIGO-BLUE

FROM

POLYGONUM TINCTORIUM AND OTHER PLANTS.

By EDWARD SCHUNCK, Ph.D., F.R.S.

(Concluded from p. 130).

Blotia Tankervillea.

THE occurrence of a blue colouring-matter in this and other plants belonging to the Orchidaceæ, such as *Calanthe veratrifolia*, was first noticed by Clamor-Marquart* and by the late Dr. Crace Calvert†. The attention of these observers was directed to these plants by seeing the blue colouration appearing in the white petals of the flowers on their beginning to fade; and they found the blue colour to be due to indigo. In accordance with the views then prevailing, they assumed the pre-existence of the colouring-matter, either as indigo-blue or as its hydride, in the tissues of these plants. It is easy to see, however, on reading the accounts of their experiments that the colouring-matter was really formed during the processes

* Buchner, Repertorium f. die Pharmacie, B. lvi. f. i.

† Journal de Pharmacie, t. vi, p. 193.

employed for its extraction; and it seemed to me, therefore, highly probable that the plant would be found to contain some glucoside similar to indican.

Bletia Tankervillea is not difficult to procure, being frequently grown for the sake of its handsome brown and white flowers and its general beauty. The leaves of the plant having been cut in pieces and ground with water between two stones to a pulp, which is strained through calico, yield a green muddy liquid which, heated to near the boiling-point, gives a thick green coagulum. The liquid filtered from this coagulum is clear, of a deep yellow colour, with a slight acid reaction and an acrid taste. When this liquid is mixed with sulphuric or hydrochloric acid and left to stand, it deposits dark-coloured flocks consisting of indigo-blue mixed with a substance which imparts a purple colour to boiling alcohol, probable indirubine. The presence of glucose may be detected in the filtrate by the usual test, while the same test applied before the addition of acid shows no indication of its presence. It may hence be inferred that the liquid contains in solution a glucoside similar to, if not identical with, indican. It undergoes, like indican, a complete change when submitted to the action of alkalis; for if its watery solution be mixed with caustic alkali and boiled, then with an excess of sulphuric acid and again boiled, it deposits brown flakes, which are found to contain but little indigo-blue, while in the filtrate only a trace of glucose can be detected. The same change takes place gradually when the watery solution is left to stand for several days at the ordinary temperature.

The gradual formation of indigo-blue in the leaves of *Bletia Tankervillea* may be easily traced in the same way as with those of *Polygonum tinctorium*, by placing the lower ends, immediately after cutting, in dilute hydrochloric acid and leaving them freely exposed for a few days. The acid, as it ascends, causes a dark discolouration; and the part discoloured, after immersion in boiling alcohol to remove the chlorophyll, appears blue.

Indigofera tinctoria.

It would be a matter of some interest to ascertain in what state the colouring-matter exists in this the most important of all the plants yielding indigo. From what I have said, however, it will be apparent that, in order to arrive at a certain conclusion, it would be necessary to work with fresh leaves; for if they contain a glucoside resembling indican, this would in a very short time undergo complete decomposition. I obtained some seeds of *Indigofera tinctoria* from Messrs. Vilmorin, Andrieux, and Co., and treated them in accordance with the directions they kindly gave me. The seeds germinated, and the young plants lived for some time in a hothouse; but unfortunately, they attained no great size, and soon decayed and died, so that I was unable to obtain a quantity of leaves sufficient for examination.

Mr. P. Michéa, an intelligent indigo-planter, with whom I have been in correspondence, gives me, however, some interesting information relating to this part of the subject. Mr. Michéa writes to me as follows:—"It was my finding in the *Indigofera* of India (in the wild plant which grows at many places in the three Presidencies as well as in the cultivated species of Bengal, the North West, and Madras) a glucoside substance perfectly similar to the indican of the *Isatis tinctoria* in all its properties, which made me declare that the colouring-matter of the Indian *Indigofera* was due to indican."

The experiments just described lead to the conclusion that in all the indigo-yielding plants hitherto examined, the colouring-matter is derived from a glucoside which splits up with great ease into indigo-blue and glucose, and that this glucoside is probably, in all cases, the same, and identical with, the indican of *Isatis tinctoria*.

Various other plants have been supposed to yield indigo-blue. Of these I have examined the following:—

Galega officinalis.

Hedysarum Onobrychis (Sainfoin).

Polygonum Fagopyrum (Buckwheat).

Polygonum Persicaria.

Rhinanthus Crista-galli.

Sophora japonica.

Spilanthes oleracea.

The leaves of these plants, when treated in the manner above described, yielded no trace of any substance resembling indican, and showed no indications of containing any blue colouring-matter like indigo.

CRYSTALLISATION OF PHOSPHORUS.

By GEORGE WHEWELL, F.I.C., F.C.S.

IN 1872, while experimenting on the action of phosphorus on nitrogen gas in sealed tubes, after melting the phosphorus and spreading it over the sides of the tube and allowing the tube to stand for several weeks, I noticed small colourless crystals begin to form. In 1874 I noticed abstracts of two papers in the *Chemical Society's Journal* (2nd series, vol. xii., p. 86g) by J. L. Smith and W. L. Herman, on the formation of crystals of phosphorus *in vacuo*. I wrote a note to the *CHEMICAL NEWS* (vol. xxx., p. 168). I did not claim priority or originality, or try in any way to detract from the merit of Messrs. Smith and Herman's papers. In Mr. Herman's letter (vol. xxx., p. 194) he tried to throw ridicule on my note, and even hinted his opinion that I had not made any crystals at all. I prepared a test-tube and sent it to Dr. Burghardt, of the Owens College, Manchester, and I append his letter, as requested by Mr. G. E. Davis (vol. xxxix., p. 115).

"The Owens College, Manchester,
"March 2, 1875.

"DEAR SIR,

"I thoroughly investigated the matter you requested me to enquire into, immediately after receiving the tube containing the phosphorus crystals. Of course you are aware that crystals of phosphorus have been obtained in other ways long ago, and it was conclusively proved that they belonged to the regular system of crystallography. The crystals in the test-tube you kindly sent me were undoubted phosphorus crystals, exhibiting the two characteristic forms observed on crystals prepared by dissolving phosphorus in rock-oil when hot and cooling, and the methods published long ago. The forms I observed were the regular octahedron and the rhombic dodecahedron, with rather rounded edges, the interfacial angles not being sufficiently well defined to allow of accurate measurements being made. There can, however, be no doubt about their being the two forms mentioned above. On opening the tube for a few seconds fumes of phosphorus came out, and the tube on being closed again, and left alone for a few days, was thickly coated with amorphous phosphorus and the crystals had vanished. . . .—Believe me, &c.,

"C. H. BURGHARDT."

"The Owens College, Manchester,
"March 18, 1879.

"DEAR SIR,

"The forms observed on the crystals were the octahedron and the rhombic dodecahedron, the former predominating; the crystals did not exhibit any sharply-defined edges or angles, consequently goniometrical measurements were not possible. The crystals were perfectly transparent. . . .—With kind regards, &c.,

"C. H. BURGHARDT."

My friend, Mr. W. H. Wood, now of the Yorkshire College, wrote me, in November, 1874:—

"I have tried your method of producing phosphorus crystals and was successful, as I could easily distinguish crystals when I examined the tube two or three days after. . . ."

Very few chemists have made crystals of phosphorus, and the only thing I claim in the above method is a very

ready means of preparing crystals of phosphorus suitable for lectures or showing it to a class when treating of phosphorus and its compounds. The method places phosphorus crystals in the hands of anyone who cares to spend ten or fifteen minutes in preparing them. I am sorry I have not studied the subject further, being fresh from College at the time, and the note being my first attempt at writing I was somewhat cowed. I came to the conclusion that Mr. Herman thought he had a vested right in crystals of phosphorus.

I have found the following the best plan for producing the crystals:—Take a piece of glass tubing and seal one end (or a test-tube); place at the bottom of the tube a piece of phosphorus about the size of two peas; draw out the other end; then heat the tube, commencing about one inch or so above the phosphorus, so as to drive out as much air as possible; then seal the open end (or cork the test-tube and make it air-tight). Melt the phosphorus, and spread it as thinly and evenly over the sides of the tube as possible. Allow to stand for a week or two, or place into a freezing mixture, when crystals will form in a few days.

END-ON ILLUMINATION IN PRIVATE SPECTROSCOPY, AND ITS APPLICATIONS TO BOTH BLOWPIPE FLAMES AND ELECTRIC ILLUMINED GAS-VACUUM TUBES.*

By PIAZZI SMYTH,
Astronomer Royal for Scotland, and past President of R.S.S. Arts.

PART I.

Defects of Former Methods, and New Conditions to be Fulfilled.

As there is nothing in the shape of splendid optical projection of grand phenomena on a large screen connected with the present paper, and as it refers merely to a practical difficulty, as met with by a private worker in a small way, in connection, too, with only a very limited branch of the general subject of spectroscopy,—some apology may be necessary for bringing so apparently trifling a matter before any public meeting. Especially, too, is such an explanation required before this long-established Society, which has had all the finer bearings of the higher spectrum analysis expounded to its members so often and so eloquently by Dr. Stevenson Macadam, and illustrated so glowingly by Mr. Hart's electric lamp, years and years ago, that not a few persons may have almost identified the subject itself, or anything about spectroscopy, with that particular and assuredly most magnificent method of demonstration. My excuse, therefore, can only consist in trying to make out, and perchance bringing home, though by mere words only, to the inner conscience of some real workers (of whom there must be many here) the exact manner of occurrence of a serious, albeit small, case of positive scientific difficulty; or in showing how a very lion of a trouble was one day found to be seated right across the special path of progress and research which I was desirous of pursuing. But how, after due interrogation as to wherefore he came to be there, the said lion was presently induced to get up, make himself scarce, and not exhibit his threatening front, to me at least, in that locality again.

If, therefore, the Society will be graciously content with such a modest bill of quiet fare as that, I will at all events endeavour to make the nature of the case very clear. With this purpose, therefore, I now commence with what is generally admitted by all the world as being foundational in spectroscopy; while it is also the necessary beginning of our subject for this evening, viz.:—

In every kind of spectroscopic observation, high or low, of fire or flame, public or private, *brilliantness* in the light operated on—so far as it can be fairly commanded—is the soul of success. In private researches, too, more particularly, where we merely look into the eye-piece of a small optic tube for our own sole and, for the time, solitary satisfaction, the mere quantity of shining matter being examined is of no sort of consequence; for we view of it at a time only as much as fills a little slit in a metal plate about $\frac{1}{16}$ inch high and $\frac{1}{32}$ inch broad. A minute, or no more than microscopic sized, particle is therefore enough, in that way; but *brilliant*, and even three times brilliant, the intrinsic light of that little particle must be, or you will be condemned to employ only the least disperse of prisms wherewith to form its spectrum; and therein, from the extreme red, all the way through the orange, yellow, green, blue, and right up to the ultimate violet and lavender grey of the spectral field, you will fail (if your spectrum is of the all important "discontinuous" order) to see anything but merely a few faint, hazy-edged, unsatisfactory bands. In place it might, or should, have been of multitudes of vivid lines, capable of almost unlimited nicety of micrometric measurement, and as firmly fixed in their several and respective spectrum places, as are the very stars of heaven in their almost eternal positions.

"Then, of course," you will say, "every sensible spectroscopist, whenever at least he is observing in a chamber upon any chemical element within his grasp, endeavours to make the illumination, or incandescence under which he looks at that particular matter, as bright as he possibly can."

Most certainly! That is exactly what he does, and even glories in doing; for he finds that he cannot make the said stuff too bright for the spectroscope, if only he preserves it at the same *temperature*. But precisely there comes in our envious and opposing lion; for almost every known attempt of man as yet to increase the brightness of a light, when at all successful in that respect, is found in the end to have enhanced its *temperature* amazingly. And if the temperature be so enhanced, where is the man who shall venture to predict, from theory alone, what the result will be in that potentially, most extensive, almost interminably long region of accurately measurable facts, the prismatic spectrum of light; wherein red is from violet, as far as north is from south?

Do you doubt there can arise any real and serious spectral changes from merely altering the temperature of one and the same flame of burning matter? Look at the metal lithium in any of its salts. How early in the history of spectroscopy did not that too little appreciated genius Fox Talbot show that chloride of lithium in a lamp flame formed one brilliant line in the scarlet-red and another very faint one in the orange-brown of the spectrum. Wherefore the brighter of them was called lithium α , the paler lithium β ; and the innocent, trusting public thought they would always know where, in the spectrum, by those letters as names, to find those lines.

But after the subsequent grand outcome of the full spectrum analysis of Professors Bunsen and Kirchhoff, and when not a few men insisted on rising from the mere beginnings of heating by flame, up to the ecstatic increase of temperature by the electric spark,—behold the lithium's red line, though still in exactly the same part of the spectrum's measured scale as before, has become woefully pale; the lithium orange line has burned up like a beacon; and far, far away in the comparatively distant regions of the spectral blue, there is a new line of almost rivaling brightness. Wherefore a new naming of the lines by Greek letters takes place among spark observers; the identical lithium β of flame-spectroscopy becoming lithium α of electric spark spectroscopy; a line hitherto unknown becoming lithium γ ; and poor old lithium α being known now only as lithium β .

But even that is not the end of the changes of brightness and name with increase of temperature; for pres-

* Read before the Royal Scottish Society of Arts, February 20, 1879.

ently still more ambitious geniuses increase the power of their galvanic batteries, the size of their induction coils, the length of their sparks, and then condense them both in time and space with Leyden-jar apparatus until the heat is something so fearful that, in the spectrum of one and the same lithium salt, it is no longer the red, nor the orange, nor even the blue line you care for now; but farther away in more distant spectral regions there burns a violet line; and beyond, yes, vastly further beyond that still, there arise exquisite lines in the ultra lavender and even grey wilds of spectral space. So then, to suit these altered brightnesses in the hotter condensed-spark spectrum, a new and yet more altered application of Greek letters, as names, takes place; and being attended to in their Memoirs by some persons, though not by others, will contribute to the future scientific confusion of many; and all on account of a mere change of temperature in a lithium light.

Or hear those first-class German observers MM. Plucker and Hittorf, at page 6 of the *Philosophical Transactions* for 1865, as to what still more extensive and positively wholesale and radical spectral changes, mere altered degrees of heat may work. "The first fact," say they, "which we discovered in operating with our tubes, guided by the above-explained principles, was the following one:—There is a certain number of elementary substances which, when differently heated, furnish two kinds of spectra of quite a different character, not having any line or band in common." So that if you have learned to recognise every line in the spectrum of, say, nitrogen at one temperature, it will not be by any of those lines, but by some perfectly different and unlikely-looking ones, that the element will manifest, or for a while conceal, itself at another temperature!

Neither does the application of a higher heat always brighten and improve a spectrum, as witness the element caesium, discovered by Professor Bunsen through means of the old flame spectrum analysis. What a refined and exquisite spectrum, too, does not caesium present in the lamp flame! So many lines it has, and all of them thin, sharp, classical almost in the orange, yellow, green; and then two gorgeous ones in the blue, but both of them so clearly defined, so admirably distinct, that I, in my small way, should have long ago established caesium as my prime standard reference in all lamp spectroscopy, if it had not been so dreadfully expensive.

Yet what becomes of this most valuable, almost invaluable, element caesium when promoted from mere flame to the higher temperature of the condensed electric spark? Alack! there is only then to be seen one faint solitary line, which has nothing in common with its departed brethren, and but little beauty of its own. With such a spectral calamity befalling poor caesium, I have been watchfully careful, in any attempted increase of the brilliancy of other spectra, to interfere as little as possible with their temperatures. And it is simply because I have recently proved the practical efficiency of a very easy method of accomplishing at least a considerable amount of brightening up, without any increase of heat whatever, in both flame and spark spectroscopy, that I have ventured now to lay the said method for merely what it may be worth, and to what extent it may be new, before the Royal Scottish Society of Arts.

(To be continued).

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Monday, March 31, 1879.

Dr. J. H. GLADSTONE, F.R.S., President, in the Chair.

This was the Anniversary Meeting of the Society.

The President presented his Annual Report on the state of the Society. The state of the Society affords

much ground for congratulation, the past year having been one of quiet prosperity. Various alterations have been made in the bye-laws, and the publication of the Society's *Journal* has been improved. 61 Fellows have been elected, and 18 have died, withdrawn, &c., during the past year, the present number of Fellows being 981. There are 34 Foreign Members and 1 Associate; 28 names are awaiting election. The Treasurer's report shows an increase of receipts over expenditure to the amount of £1964 9s. 5d. (including a legacy of £1000 from the late Mr. Sidney Ellis). 68 papers have been read before the Society during the past year. Lectures have been given by H. C. Sorby and S. H. Vines. The number of papers from 1859 to 1869 averaged 36 annually, fell as low as 22 in 1872, and since has rapidly increased. There is an increasing disposition to illustrate papers by experiments: this practice adds much to the interest of the meetings of the Society. On the 13th of November last Prof. Würtz delivered the Faraday Lecture at the Royal Institution, "Sur la Constitution de la Matière à l'état Gazeux." Those who were present will not readily forget the perfect lucidity, the rare manipulative skill, or the enthusiasm of the Lecturer on that occasion. The Council have carefully considered the condition of the Library and its rules. It has been resolved that no serial publication of which the Society does not possess a duplicate copy shall be taken away from the rooms. Duplicate copies of back numbers of *Liebig's Annalen* and the *Annales de Chimie et de Physique* have been ordered, and these works, together with other important serial publications, are now being taken in in duplicate. Many new books have also been ordered, and on this point the Council would be glad of suggestions from Fellows of the Society. Arrangements have been made for Fellows at a distance to have books sent to them on giving a written order and paying the cost of carriage. A new series of instructions has been drawn up for the use of abstractors. Great efforts have been made to publish the *Journal* within the first week of each month. The ventilation of the meeting-room has been much improved by the combined investigations of Mr. Perkin and Dr. Russell, and alterations suggested by them. A new plan of choosing the President has been adopted this year. As the Faraday Lecture occurs only once every three years, whilst the President is elected for two years, it was suggested that it would be better to elect some Fellow who has already filled the office of President for one year, so that the Faraday Lecture would occur in the second year of the next President's tenure of office. A separate Report of the Research Fund is submitted, the President suggesting that a portion might, with peculiar fitness, be devoted to the accurate determination of chemical constants. A short biography of each of the Fellows deceased was then read, viz., of M. Malaguti (Foreign Member) Messrs. W. Baker, A. Bird, W. A. Lyttle, W. A. Stewart, and J. Wiggan. The President concluded his address as follows:—"While there is much to encourage us in the progress of the past twelve months, I think I only express the general feeling when I say that we ought not to feel satisfied with present attainments. The Society exists 'for the general advancement of Chemical Science;' this means both the encouragement of research and the diffusion of the knowledge of new discoveries. With regard to the promotion of research, as our members now exceed 1000, and the laboratories of our land are growing in importance, we may surely look for a larger amount of original work in coming years: we should also seek, not only to add to previous knowledge, but to increase what I may venture to term the scientific culture of the workers. In our laboratories we are isolated, and are apt to look upon our own pet subject as of prime importance; but when we meet in these rooms, or turn over the pages of our *Journal*, we are carried away to many different fields of thought in succession. This promotes largeness of view, and must react favourably upon the cultivation of our individual corner of the great field. I trust that our Society will never devote its energies too exclusively to

one branch of our science, but will lay claim to its fair share of those borderlands in which the work of the chemist blends with that of the physicist, the geologist, and the physiologist. Scientific culture will also lead to a perception of what are the higher aims of chemical enquiry. The formation of new compounds is valuable, but we are liable to be encumbered by the richness and quantity of our materials, and to forget the necessity of grouping them together as a part of systematic knowledge. The construction of the most expressive formulæ is useful, but we must always be ready to modify these as the exigencies of further knowledge may require. We want, also, to know more of the chemical force itself, and how it acts; we want to distinguish those properties which are so profoundly connected with the ultimate molecules of matter that they are little, if at all, affected by chemical combination; from those which are the sport of every change; we want to study all the transformations of energy involved in the phenomena of dynamical chemistry, and to determine with precision how the chemical force is related to the other great forces of nature. As to the diffusion of chemical knowledge, our *Journal* is the main instrument in our hands for effecting this. But the new arrangements in regard to the lending of books from the Library will doubtless advance the same object. It is happily the case that a knowledge of chemistry is fast finding its way into our upper and middle class schools; and though our science is not recognised by the Government code, many attempts are being made to introduce into our elementary schools some primary knowledge of those facts and principles of nature which lie at the foundation of chemical and physical science. Technical education is also rising into favour, and the formation of a technical college is now engaging the attention of the great City companies. The Society, as such, can perhaps do but little in this direction, although the practical applications of chemistry are directly alluded to in our Charter, but its individual members may accomplish much.

Dr. ARMSTRONG then read a list of grants from the Research Fund made during the past year:—£50 to Mr. Hartley for an Investigation of the Absorption of the Ultra-violet Rays of the Spectrum by Organic Substances; £30 to Dr. W. Ramsay for Determining the Electric Conductivity and Resistance of Solutions of Salts at Different Temperatures; £50 to Dr. Tilden for an Investigation into the Chemical Nature of the Terpenes; £10 to Mr. Shenstone for an Examination of Certain Reactions of Brucine and Strychnine; £20 to Mr. W. Jago for a Research on the Organic Matter in Sea Water; £20 to Mr. Francis Jones for the Investigation of Boron Hydride; £15 to Mr. F. D. Brown for the Experimental Study of the Theory of Fractional Distillation; £10 to Dr. Burghardt for the Investigation of the Constitution of Topaz; £15 to Prof. Thorpe for the Investigation of Abietine; £30 to Dr. Dupré for the Estimation of the Organic Carbon in Atmospheric Air. The following gentlemen who have received grants have communicated papers to the Society:—Messrs. Johnson, Carleton Williams, Drs. Harrow and Wright; Mr. Hartley in conjunction with Mr. Hartington has presented his first results to the Royal Society, *Proceedings*, xxviii., 233. The following gentlemen, who, for various reasons, have not yet communicated papers, have sent in reports to the Committee:—Drs. Crow, Carnelly, Tilden, and Ramsay; Messrs. Bedson, Neison, Shenstone, and Jago.

Dr. ODLING then moved a vote of thanks to the President and the adoption of his report. It was very gratifying to hear such a flourishing report, and thought the Fellows might congratulate themselves on having had such a President.

Mr. NEISON, in seconding the motion, urged that a copy of the General Index should be presented to every Fellow of the Society.

Dr. WRIGHT suggested that the hours during which the

library was open in the evening should be extended from seven to nine to seven to ten.

The motion was then carried by acclamation.

The PRESIDENT felt greatly honoured by the cordial manner in which the motion had been received and the kind way in which Dr. Odling had introduced it. He had experienced much pleasure during his occupation of the chair, which any chemist would be proud to occupy. He was very grateful for the way in which the Council and the Society had seconded him in everything.

The TREASURER (Dr. Russell) then read his account of the finances of the Society. The state of the Society was very satisfactory. The income for the year was £2350; the expenditure £2300. The balance in hand of the Research Fund amounted to about £230.

Mr. FRISWELL then read the report of the Auditors, Messrs. Spiller, Thomson, and Friswell.

Dr. THUDICHUM had listened with great satisfaction to the statement of the Treasurer. After making a few remarks as to the present method of electing Fellows, Dr. Thudichum proposed a cordial vote of thanks to the Treasurer, who had performed his important duties in the most perfect manner the Society could wish.

Dr. GILBERT seconded the motion, which was carried unanimously, with much pleasure.

Dr. RUSSELL returned thanks to the meeting.

Mr. NEISON proposed a vote of thanks to the officers and Council.

This was seconded by Mr. GROSJEAN, and carried unanimously.

Mr. PERKIN replied.

Votes of thanks were subsequently given to the Auditors, the Editors, Abstractors, and the Reporter of the Society.

The following officers were then announced from the chair as having been duly elected for the ensuing year:—President, Warren De La Rue, F.R.S.; Vice-Presidents, F. A. Abel, C.B., Sir B. C. Brodie, E. Frankland, J. H. Gladstone, A. W. Hofmann, W. Odling, Lyon Playfair, A. W. Williamson, F. Field, J. N. Gilbert, N. S. Maskelyne, H. E. Roscoe, R. Angus Smith, J. Young; Secretaries, W. H. Perkin, H. E. Armstrong; Foreign Secretary, Hugo Müller; Treasurer, W. J. Russell; other Members of the Council, M. Carteighe, A. H. Church, W. N. Hartley, C. W. Heaton, E. Riley, W. C. Roberts, W. A. Tilden, W. Thorpe, T. E. Thorpe, J. T. W. Thudichum, R. V. Tuson, R. Warington.

PHILOSOPHICAL SOCIETY OF GLASGOW.

CHEMICAL SECTION.

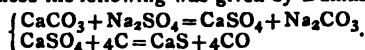
Ordinary Meetings.

Mr. JAMES MACTEAR, President, in the Chair.

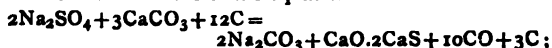
"On the Manufacture of Phosphorus," by Mr. JAMES READMAN. The author remarked at the outset that his communication contained nothing absolutely new, but was simply an account of the practical aspects of the phosphorus manufacture. Various mineral phosphates are now used in the manufacture of phosphorus; bone-ash is no longer remunerative, on account of its high price. Among the varieties of mineral phosphate are—Canadian phosphate, German or Nassau phosphate, Charleston phosphate, and Sombbrero phosphate. The first stage in the manufacture of phosphorus is to decompose the calcium phosphate completely in a large cylinder with sulphuric acid of 110° to 115° Twaddle, with constant agitation. The calcium sulphate is then filtered off, and the filtrate is evaporated to about 80° or 90° T., and then allowed to cool. It contains over 25 per cent of P_2O_5 . It is then mixed with coarse wood-charcoal, and dried in a muffle-furnace. The proportion of charcoal to liquor is 1 to 5. This substance contains the phosphoric acid in a partially insoluble state, so that it is different in its properties from

meta-phosphoric acid. The mixture is then transferred to retorts of Stourbridge clay, capable of holding 30 to 40 lbs. The malleable iron pipe through which the phosphorus distils is then luted on, and the heat is raised to bright redness. The phosphorus distils over, and is condensed in water. It only remains to cast it in moulds, when it is sent to market. Mr. Readman, in contradiction to the usual assertions in text-books, called especial attention to the fact that mono-calcium phosphate is not used as a source of phosphorus, for the calcium takes up valuable room, and the compound requires a much more intense heat to effect its decomposition. Redonda phosphate of aluminium is regarded by the author as the future source of phosphorus, but as yet no attempt to procure phosphorus from it has been peculiarly successful.

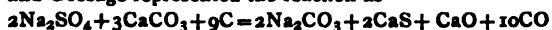
The PRESIDENT read a paper on the "*Leblanc Soda Process*." He gave an historical *resumé* of the process of alkali manufacture, mentioning that before Leblanc brought out his process it was very probably carried on as a secret process in England. The various reactions imagined by different investigators to take place were then alluded to. Among these the following was given by Dumas:—



It was afterwards supposed that an oxysulphide of calcium was formed in virtue of the equation—



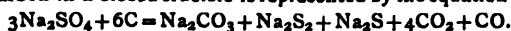
and Gossage represented the reaction as—



The author then gave an account of his own researches on the subject. As it is impossible to arrive at any definite conclusion by operating on the large scale, for different samples of black-ash, even when drawn with the greatest possible care from the same charge, show the most varied composition, the reaction was simplified, sodium sulphate and carbon being heated together in small quantity: the proportions and temperature were varied in each experiment. Some preliminary experiments were made, and proved that sodium carbonate is formed, especially at a red heat. At a higher temperature very little Na_2CO_3 was produced. The reaction evidently approximated to—



At 1250° F. the reaction between sodium sulphate and carbon in a closed crucible is represented by the equation—



In an open crucible the excess of sulphur is burned off, and it is probable that if more carbon were added all the sulphate of sodium would be converted into carbonate. On raising the temperature the Na_2S_2 and Na_2CO_3 are decomposed, and the only products are $\text{Na}_2\text{S} + \text{CO}$.

The next paper was read by Mr. COLEMAN. Its subject was "*The Liquefaction of Gases*." The author gave a description of his apparatus for liquefying hydrocarbons with low boiling-points obtained in the distillation of shale, and explained it on theoretical principles.

NOTICES OF BOOKS.

The Science Index: a Monthly Guide to the Contents of the Scientific Periodicals. January, 1879. Vol. I., No. 1.

We were very much pleased with the fundamental idea of this journal as conveyed in its title, but on looking further we experienced no small degree of disappointment. A relatively trifling portion of the space is devoted to science, whilst the lion's share is allotted to the industrial arts and to commerce. Thus, of the 112 columns, chemistry receives 3; astronomy, $1\frac{1}{2}$; electricity, $1\frac{1}{2}$; natural history, 1; physics, 2; geology, $1\frac{1}{2}$; botany, $\frac{3}{4}$; math-

matics, $\frac{1}{2}$; whilst physiology and psychology are absent altogether. On the other hand, architecture is favoured with 8 columns, commerce with 4, capital and labour with the same space, lawsuits with 2, and railways with 4. Hence we submit that the new periodical is altogether wrongly named. We observe, further, that the French, German, and other Continental scientific journals do not appear to be indexed. Amongst our American contemporaries the *Scientific American* with its supplement is alone selected, to the exclusion of not a few journals of high and recognised value. Many British periodicals are also overlooked. Thus we see no mention of the *Chemist and Druggist*, the *Pharmaceutical Journal*, the *Analyst*, the *Sugarcane*, &c. The medical journals are omitted in a body, as are also several important papers devoted to various branches of natural history. Now it must be clear that a partial and incomplete index is of very doubtful value. A student hears a rumour concerning a certain memoir which has appeared. He searches the "Science Index," and finding no mention of it is led to conclude that it does not exist, the fact being that it is to be found in some journal which has here been overlooked.

Another defect is that certain memoirs are credited not to the journals in which they first appeared, but to others in which they have been reproduced. Typographical errors are also too numerous, and in certain cases are of a misleading character.

If the editor in future issues will attend to these suggestions he may achieve a decided success. But we must confess ourselves unable to recommend the January number as a "Science Index."

The Textile Colourist. Vol. I., No. 1.

This is a new periodical devoted to dyeing, bleaching, printing, &c., and has just appeared at Philadelphia under the management of Dr. Frank. It contains a number of practical recipes illustrated by well dyed samples of dyed yarns, trimmings, &c. Concerning the chapter on mordants, by M. P. Prunier, we will merely remark that he gives the name "iron liquor," not as usual to the acetate of iron, but to a nitro-sulphate, and that he terms his perchloride of tin "red liquor," a name formerly peculiar to the acetate of alumina. This, we fear, will lead to further confusion in a nomenclature which is already complicated enough.

We wish our new contemporary a prosperous and a useful career.

CORRESPONDENCE.

MANUFACTURE OF SULPHURIC ACID.

To the Editor of the Chemical News.

SIR,—Since in your review of a new treatise on the manufacture of sulphuric acid my authority is referred to on a special point, that of the introduction of nitric acid in the liquid form, I feel bound to state that I am no longer so decidedly in favour of that system as I certainly was some years ago. Having investigated this matter very carefully, along with all other points of sulphuric acid and alkali making, on visits to the prominent English, German, Austrian, and French Alkali works, I have come to the conclusion that, *on the whole*, the decomposition of the nitrate and sulphuric acid by the heat of the burner gas, as practised in England and at many Continental works, is the better plan, *if carried out in the proper way*. In that case the consumption of nitrate is just as low as when employing nitric acid in the liquid form. The difference between the two plans, as far as cost and convenience are concerned, is certainly very slight now, since the nitric acid is now no longer introduced by "cascades"

but it is, by some of the most careful German and Austrian manufacturers, simply run down the Glover tower, all fear of any appreciable loss of nitrous compounds in that tower having vanished. I cannot, of course, explain these matters in detail here; this is done in my "Treatise on the Alkali Manufacture," the first volume of which (Sulphuric Acid) has just been published in German, whilst the English edition (published by Mr. John van Voorst) is far on its way through the press.—I am, &c.,

GEORGE LUNGE.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 10, March 10, 1879.

This issue is taken up with the annual presidential address and with an account of the awards of the various prizes offered for certain branches of research. One of these reports, relative to the prize awarded to M. Turpin for his non-poisonous pigments, requires notice. These, we are told, include not merely certain colours long known, such as zinc-white and chromate of zinc (I) but "others which are new and absolutely inoffensive, as they derive their origin from coal-tar." Is not this a somewhat hazardous conclusion?

Bulletin de la Société Chimique de Paris,
No. 3, February 3, 1879.

Researches on Strychnine.—H. Gal and A. Etard.—The authors, on treating strychnine with hydrate of baryta, have obtained two new hydrated bases, di- and trihydro-strychnine.

On Succinic Fermentation.—P. Miquel.—The author has detected the bacterium which effects the decomposition of asparagin into succinate of ammonia. It is a mobile organism formed of one, two, or rarely of a greater number of joints. The mature being and its germs are destroyed by a temperature of 48° to 49° kept up for two hours.

Dextro-rotatory Amylic Alcohol.—J. A. Le Bel.—Lævo-rotatory amylic alcohol, indicating -4.33° for 10 centimetres of a liquid column may be rendered inactive by its transformation into amylate of sodium and by the action of an elevated temperature upon the latter. The author separates the dextro-rotatory alcohol from the inactive kind by destroying a portion of the lævo-rotatory kind by moulds.

Action of Diastase, Saliva, and Pancreatic Liquid upon Starch and Glycogen.—F. Musculus and J. de Méring.—Already noticed.

Aniline and the Methylated Toluydins, and on their Coloured Derivatives.—P. Monnet, F. Reverdin, and E. Noelting.—The authors describe mono-methyl-aniline, dimethyl-aniline, mono-methyl-ortho-toluydin, dimethyl-ortho-toluydin, mono-methyl-meta-toluydin, dimethyl-meta-toluydin, and dimethyl-para-toluydin. It results from their researches that among all the bases found in commercial methyl-aniline pure dimethyl-aniline itself is the only one which can be advantageously used for the manufacture of violet. Mono-methyl-aniline and dimethyl-ortho-toluydin give, indeed, good violets of a redder tone than that of dimethyl-aniline, but the yield is very trifling. Mono-methyl-ortho-toluydin gives a good return of violet, but it has the defect of being insoluble in water. The methylated derivatives of the two other toluydins yield brown and grey colours of no value.

Gazzetta Chimica Italiana.

Anno viii., 1878. Fasc. viii. and ix.

Cumo-phenol-carbonic Acid.—E. Paterno and G. Mazzara.—This acid takes the form of flat needles or nacreous laminæ, which melt at 120.5° and volatilise without decomposition; sparingly soluble in cold water, more so in hot water, and very soluble in alcohol and ether. The aqueous solution yields with ferric salts a very intense violet-blue colouration. Its composition is expressed by the formula $C_{10}H_{12}O_3$. The authors have examined its barium, lead, and silver salts.

Alleged Existence of Oxygenated Water in the Organism of Plants.—Prof. Giuseppe Belucci.—The presence of hydric peroxide in the juices of plants was first admitted by Schœnbein, and was maintained more recently by Clermont. The assumption is refuted by the author's experiments.

Two Propyl phenols and other Derivatives of Propyl-benzol.—Dr. P. Spica.—The author describes synthetic propyl-benzol obtained by the method of Fittig, Schäffer, and König, its sulph-acids and the α - and β -derivatives of the latter.

Dimorphism of the Aceto-toluides.—Dr. R. Panebianco.—An optical and crystallographical study, not admitting of useful abstraction.

Synthesis of Phenyl-cinnamic Acid.—Dr. A. Ogliarolo.—The author obtains the new acid by heating in a reflux apparatus for eight hours, at a temperature of 150° to 160° , about 25 grms. of phenyl-acetate of sodium, previously dried at 110° to 120° , with 16 grms. of benzoic aldehyd and 60 of anhydrous acetic acid. The product of the reaction, which, when hot, is entirely liquid, condenses on cooling to a reddish brown crystalline mass, which is diluted with water and boiled to expel the excess of acetic acid employed. The result is filtered when the filtrate deposits the acid in white needles.

Preparation of Ammonialdehyds with Mixed Radicles.—R. Schiff.—An examination of the action of benzoic aldehyd upon chloral-ammonium and butyl-chloral-ammonium.

Studies on Teucrium Fruticans.—Dr. A. Ogliarolo.—The author has obtained from this plant a compound which he provisionally names teuclin, and has examined its behaviour with nitric and sulphuric acids.

Chemical Nature of the Essence of Laurocerasus and of Bitter Almonds.—Dr. M. Fileti.—The author has studied the action of nascent hydrogen upon this essence and upon amygdalin.

The Alkaline Polysulphides as Reagents for Cobalt.—G. Papasogli.—Among the characteristic reactions of cobalt is one founded upon the blood-red colouration (or rose, in small quantities) obtained if there is poured into the solution of the double cyanide of cobalt and potassium some nitrite of potassium with nitric acid. A similar colouration is obtained if, instead of adding nitrite of potassium and nitric acid to the above-mentioned cobalt solution, there is added a little drop of a solution of a yellow alkaline sulphide. In order that the reaction may be most sensitive the polysulphide should be added in such a manner that the two liquids may not completely mix but form two distinct strata. If cobalt is present the plane of separation will show a blood-red, more or less intense according to the quantity of the metal. The author has obtained this reaction with $\frac{1}{2}$ c.c. of a liquid containing 0.0005 grm. of cobalt. The presence of nickel does not interfere.

Glucoside of Liquorice.—Prof. F. Sestini.—Not susceptible of useful abstraction.

Chemiker Zeitung.

No. 9, 1879.

Sugared Ultramarine.—C. Fürstenau.—The Austrian ultramarine manufacturers sophisticate their colours with

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

FEBRUARY, 1879.

The following are the returns of the Society of Medical Officers of Health:—

	Appearance in a foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	An- Sulphuric hydride.	Hardness on Clark's Scale.	
		Saline.	Organic.								Before Boiling.	After Boiling.
<i>Thames Water Companies.</i>												
Grand Junction	Clear	0'000	0'007	0'180	0'061	22'10	8'730	0'720	1'08	1'610	14'3	4'60
West Middlesex	Clear	0'000	0'009	0'180	0'047	22'00	8'890	0'684	1'15	1'430	13'2	4'60
Southwark and Vauxhall	Clear	0'000	0'008	0'150	0'036	21'70	8'460	0'720	1'15	1'780	13'2	3'30
Chelsea	Clear	0'000	0'007	0'135	0'062	22'20	7'770	0'579	1'15	1'200	13'7	3'70
Lambeth	Clear	0'000	0'006	0'135	0'087	21'70	6'440	0'710	1'29	1'740	13'2	3'30
<i>Other Companies.</i>												
Kent	Clear	0'000	0'003	0'360	0'002	30'90	8'490	1'222	1'80	3'000	18'8	6'50
New River	Clear	0'000	0'006	0'180	0'032	22'40	7'280	0'540	1'15	1'160	14'3	3'70
East London	Clear	0'000	0'007	0'180	0'062	26'00	9'570	0'600	1'36	1'710	16'5	4'60

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours; and in the case of the metropolitan waters the quantity of organic matter is about eight times the amount of oxygen required by it.

C. MEYMOTT TIDY, M.B.

3 parts of gypsum, and lest the colour should appear too pale it is further mixed with glycerin or glucose, or a mixture of both. This keeps the powder damp and renders the colour apparently deeper.

Formation of Ammonium Nitrite.—V. Löseke.—When water evaporates in air a little ammonium nitrite is always formed. The author shows that this takes place at the expense of the free nitrogen of the atmosphere, more of which enters into combination at low than at high temperatures.

Industrial Activity.—At a general meeting of the German Distillers' Association it was resolved to establish and maintain a chemical laboratory, an experimental distillery, a school of distilling, a trade journal, a glass-blowing establishment (for the manufacture of normal hydrometers), and an office for general intelligence.

No. 10, 1879.

Chloride of Lime.—Dr. Filsinger.—The author states that in case of English samples the importers have very frequently to complain of differences in strength, whilst in German qualities important deviations from the stipulated strength are almost unknown.

Spurious Seeds.—According to Dr. A. Stutzer the manufacture of artificial clover seed is now a flourishing business in Germany. Fragments of gravel of a suitable size are obtained by sifting, and are then agitated with certain colouring matters in a revolving drum till their appearance is considered satisfactory.

Unwholesome Honey.—A sample of honey which had occasioned illness on consumption was found when microscopically examined to contain multitudes of Acari (*Glyciphagno prunorum* and *agilis*).

Manufacture of Picric Acid.—J. Marzelli proposes to add slowly the sulphacid of phenol to concentrated nitric acid. The reaction proceeds slowly without requiring the application of heat. The crystals of crude picric acid are washed in cold water, pressed, and recrystallised.—*Monit. Prod. Chimiques*.

Cerium Aniline-Black.—J. Lytsche, of St. Petersburg, has used this process successfully for upwards of a year, employing a cerous sulphate formed by dissolving the cerite of Riddarhytta, in Sweden, in sulphuric acid. The black is said to be faster than that produced with vanadium.—*Engl. Polyt. Journ.*

No. 11, 1879.

Utilisation of Animal Refuse.—Dr. B. Terne.—The author points out that offensive gases are invariably generated when fresh animal matters are boiled by means of steam at a high pressure, and recommends that they should be gradually passed through the furnace.

Sensitiveness of Chemical Balances.—A. Verbeek.—The author, by his compensating arrangement of the axial knife-edges of the balance, renders it equally sensitive when loaded as when empty.

Saccharification of Starch.—M. Riban.—If starch is soaked for a year in a cold saturated solution of common salt it is gradually converted into glucose.

Reaction of Bile Pigments in Urine.—About 2 grms of the urine are poured into a test-tube, acidulated with 2 or 3 drops of concentrated sulphuric acid, and a small crystal of potassic nitrite is introduced, taking care that it does not fall upon the side of the tube. If bile pigments are present splendid grass-green stripes are immediately formed in the liquid.—*Repertoire de Pharmacie*, 35, 58.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 10, March 6, 1879.

M. Adam lays great emphasis on the use of soot in horticulture and agriculture as a means of destroying or banishing noxious insects.

MISCELLANEOUS.

Russian Scientific News.—Some experiments executed in Cronstadt, have shown that an explosion of two grammes of nitro-mannite, placed into a capsule, produces a full detonation of damp gun-cotton, containing at least 25 per cent of water. The explosion of the nitro-mannite was effected by the explosion of 0'2 grm. of mercury fulminate, and in other experiments by using the same amount of diazo benzol nitrate. Other fulminating compounds (KClO_3 , $\text{C}_6\text{H}_2(\text{NO}_2)_3$, OK) gave the same effect, but the quantity of them, wanted for the same purpose, exceeds 1 grm.

M. Adrianovsky reports on the action of aluminium chloride on acetic and sulphurous anhydrides. Acting by aluminium chloride on acetic anhydride at ordinary tem-

perature acetyl chloride and aluminium acetate are produced. Sulphurous anhydride with Al_2Cl_6 gives a compound having the following composition:— $AlCl_2SO_2Cl$. This reaction takes place at ordinary temperature, but is more easily conducted at 50° to 60° Celsius.

Prof. Bexetoff has made some new experiments on the absorption of hydrogen by palladium, and determined the atomic heat of the hydrogen when alloyed with palladium. The atomic heat was found to be 5.88. 25.0938 grms. of palladium in the form of wire and strips were used for the experiment. The metal absorbed 0.1418 grm. of hydrogen, corresponding to 1575 c.c.; in other words, the palladium absorbed 710 volumes of the hydrogen.

SERGIUS KERN, M.E., St. Petersburg

NOTES AND QUERIES.

Testing of Coal-Tar Products.—Can any of your readers recommend a book on the commercial testing of coal-tar products?—T. C. W.

Gases from Vitriol Chambers.—(Reply to W. L.)—You will find the information you desire in our new work on sulphuric acid, published by Sampson, Low, and Co., 188, Fleet Street, E.C., and noticed in last week's CHEMICAL NEWS.—LOCK AND LOCK.

MEETINGS FOR THE WEEK.

- MONDAY, 7th.—Medical, 8.30.
— Royal Institution, 5. General Monthly Meeting.
TUESDAY, 8th.—Civil Engineers, 8.
— Photographic, 8.
— Anthropological, 8.
WEDNESDAY, 9th.—Geological, 8.
— Microscopical, 8.

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In consequence of GOOD FRIDAY occurring in the ensuing week, the CHEMICAL NEWS will be published on Thursday next, April the 10th. Advertisements must therefore be forwarded to the Office not later than 2 o'clock on Wednesday, the 9th instant.

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INSTITUTE OF CHEMISTRY.

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 Fellows of the Institute who shall before the 31st December next have
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 scribed practical examination, and successful competition for these
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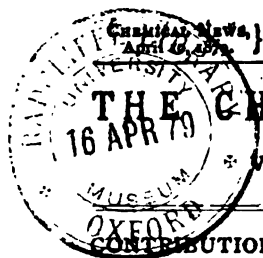
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THE CHEMICAL NEWS.

Vol. XXXIX. No. 1011.

CONTRIBUTIONS TO MOLECULAR PHYSICS IN HIGH VACUA.

By WILLIAM CROOKES, F.R.S.

THIS Paper is a continuation of one "On the Illumination of Lines of Molecular Pressure, and the Trajectory of Molecules," which was read before the Royal Society on the 5th of December last. The author has further examined the action of the molecular rays electrically projected from the negative pole in very highly exhausted tubes, and finds that the green phosphorescence of the glass (by means of which the presence of the molecular rays is manifested) does not take place close to the negative pole. Within the dark space there is absolutely no phosphorescence; at very high exhaustions the luminous boundary of the dark spark disappears, and now the phosphorescence extends all over the sensitive surface. Assuming that the phosphorescence is due either directly or indirectly to the impact of the molecules on the phosphorescent surface, it is reasonable to suppose that a certain velocity is required to produce the effect. The author adduces arguments to show that within the dark space, at a moderate exhaustion, the velocity does not accumulate to a sufficient extent to produce phosphorescence, but at higher exhaustions the mean free path is long enough to allow the molecules to get up sufficient speed to excite phosphorescence. At a very high exhaustion there are fewer collisions, and the initial speed of the molecules close to the negative pole not being thereby reduced, phosphorescence takes place close to the pole.

Experiments are described in which a pole folded into corrugations is used at one end of a tube, the pole at the other end being flat set obliquely to the axis of the tube, and having a plate of mica in front pierced with a hole opposite the centre of the pole. The questions which this apparatus was designed to answer are:—(1.) Will there be two sets of molecular projections from the corrugated pole when made negative, one perpendicular to each facet, or will the projection be perpendicular to the electrode as a whole, *i.e.*, along the axis of the tube? (2.) Will the molecular rays from the oblique flat pole, when this is made negative, issue through the aperture of the screen along the axis of the tube, *i.e.*, direct to the positive pole, or will they leave the pole normal to the surface and strike the glass on its side? With the corrugated pole experiment shows that at high exhaustions molecular rays are projected from each facet to the inner surface of the tube, where they excite phosphorescence, and form portions of ellipses by the intersection of the planes of molecular rays with the cylindrical tube. When the oblique flat pole is made negative, a stream of molecules shoots from it nearly normal to its surface, and those which pass through the hole in the plate of mica strike the side of the tube, forming an oval patch of a green colour.

The oval patch in this apparatus happens to fall on a portion of the glass which has previously had its phosphorescence excited by the molecular discharge from the other corrugated pole. The phosphorescence from this pole is always more intense than that from the flat pole, and the glass, after having been excited by the energetic bombardment, ceases to respond readily to the more feeble excitement from the flat pole. The effect, therefore, is, that when the oval spot appears, it has a dark band across it where the phosphorescence from the other pole had been

taking place. The glass recovers its phosphorescent power to some extent after rest.

In this apparatus a shifting of the line of molecular discharge is noticed. If the coil is stopped and then set going repeatedly, always keeping the oblique pole negative, the spot of green light occurs on the glass at the spot where it should come supposing the discharge were normal to the surface of the pole. But if once the flat pole is made positive, the next time it is made negative the spot of light appears nearer the axis of the tube, and instantly shifts to its normal position, where it remains so long as its pole is made negative. There seems no limit to the number of times this experiment can be repeated.

A suggestion having been made by Professor Stokes that a third, idle, pole should be introduced between the negative and positive electrodes, experiments are described with an apparatus constructed accordingly. The potential of the idle poles (of which there are two) at low exhaustions is very feebly positive; as the exhaustion gets better the positive potential increases, and at a vacuum so good as to be almost non-conducting, the positive potential of the idle poles is at its greatest. The result is that an idle pole in the direct line of fire between the positive and negative poles, and consequently receiving the full impact of the molecules driven from the negative pole, has a strong positive potential.

It is found that when the shadow of an idle pole is projected on a phosphorescent screen the trajectory of the molecules suffers deflection when the idle pole is suddenly uninsulated by connecting it with earth. The same result is produced by connecting the idle pole with the negative wire through a very high resistance, such as a piece of wet string, instead of connecting it with earth. A tube, which has already been described in a paper read before the Royal Society on December 5th last, is used to illustrate this deflection. The shadow of an aluminium star is projected on a phosphorescent screen. So long as the metal star is insulated the shadow remains sharp, but on uninsulating the star by connecting it with an earth wire the shadow widens out, forming a tolerably well-defined penumbra outside the original shadow, which can still be seen unchanged in size and intensity. On removing the earth connexion the penumbra disappears, the umbra remaining as before.

It is also found that the shadow of the star is sharply projected when it is made the positive pole, the negative pole remaining unchanged.

These experiments are explained by the results just mentioned, that the idle pole, the shadow of which is cast by the negative pole, has strong positive potential. The stream of molecules must be assumed to have negative potential; when they actually strike the idle pole they are arrested, but those which graze the edge are attracted inwards by the positive potential and form the umbra. When the idle pole is connected with earth, its potential would become zero were the discharge to cease; but inasmuch as a constant supply of positive electricity is kept up from the passage of the current, we must assume that the potential of the idle pole is still sufficient to more than neutralise the negative charge which the impinging molecules would give it. The effect, therefore, of alternately uninsulating and insulating the idle pole is to vary its positive potential between considerable limits, and consequently its attractive action on the negative molecules which graze its edge. The result is a wide or a narrow shadow, according to circumstances.

After a definite shadow is produced, it is found that increasing the exhaustion makes very little change in the umbra, but it causes the penumbra to increase greatly in size. Experiments recorded in the paper already quoted have proved that the velocity of the molecules is greater as the vacuum gets higher, and consequently the trajectory of the molecules under deflecting action, whether of a magnet or of an insulated idle pole, is flatter at high than at low vacua.

An experiment is next described, having for its object to

ascertain whether two parallel molecular rays from two adjacent negative poles attract or repel each other. It is considered that if the stream carries an electric current, attraction should ensue, but if they are simply streams of similarly electrified bodies, the result would be repulsion. Experiment proves that the latter alternative happens, lateral repulsion taking place between two streams moving in the same direction.

Many experiments are given to illustrate the law of action of magnets on the molecular stream, but the results are of too complicated a character to bear condensation without the diagrams accompanying the original paper.

The molecular stream is sufficiently sensitive to show appreciable deflection by the magnetism of the earth.

The author, after numerous experiments, has succeeded in obtaining continuous rotation of the molecular stream under the influence of a magnet, analogous to the well-known rotation at lower exhaustions. Comparative experiments are given with a "high vacuum" tube, where no luminous gas is visible, but only green phosphorescence on the surface of the glass, and a "low vacuum" tube in which the induction spark passes in the form of a luminous band of light joining the two poles. These two tubes are mounted over similar electro-magnets, the direction of discharge being in a line with the axis of the magnet. Numerous experiments, the details of which are given in the paper, show that the law is not the same at high as at low exhaustions. At high exhaustions the magnet causes the molecular rays to rotate in the same direction, whether they are coming towards the magnet or going from it; the direction of rotation being entirely governed by the magnetic pole presented to the stream. The north pole rotates the molecular discharge in a direct* sense, independent of the direction in which the induction current passes. The direction of rotation impressed on the molecules by a magnetic pole is opposite to the direction of the electric current circulating round the magnet. These results offer an additional proof that the stream of molecules driven from the negative pole in high vacua do not carry an electric current in the ordinary sense of the term.

The author, after giving details of experiments in which platinum and glass are fused in the focus of converging molecular rays projected from a concave pole, describes observations with the spectroscopic, which show that glass obstinately retains at even a red heat a compound of hydrogen—probably water—which is only driven completely off by actual fusion.

The permanent deadening of the phosphorescence of glass is shown by projecting the shadow of a metal cross on the end of a bulb for a considerable time. On suddenly removing the cross, its image remains visible, bright upon a dark ground.

One of the most striking of the phenomena attending this research is the remarkable power which the molecular rays in a high vacuum have of causing phosphorescence in bodies on which they fall. Substances known to be phosphorescent under ordinary circumstances shine with great splendour when subjected to the negative discharge in a high vacuum. Thus Becquerel's luminous sulphide of calcium has been found invaluable in this research for the preparation of phosphorescent screens whereon to trace the paths and trajectories of the molecules. It shines with a bright blue-violet light, and when on a surface of several square inches is sufficient to faintly light a room.

The only body which the author has yet met with which surpasses the luminous sulphides, both in brilliancy and variety of colour, is the diamond. Most diamonds from South Africa phosphoresce with a blue light. Diamonds from other localities shine with different colours, such as bright blue, apricot, pale blue, red, yellowish green, orange, and pale green. One very beautiful diamond in the author's collection gives almost as much light as a candle when phosphorescing in a good vacuum.

Next to the diamond, alumina and its compounds are

the most strikingly phosphorescent. The ruby glows with a rich full red, and it is of little consequence what degree of colour the stone possesses naturally, the colour of the phosphorescence is nearly the same in all cases; chemically prepared and strongly ignited alumina phosphoresces with as rich a red glow as the ruby. The phosphorescent glow does not therefore depend on the colouring-matter. E. Becquerel* has shown by experiments with his phosphoroscope that alumina and many of its compounds phosphoresce of a red colour after insolation.

Nothing can be more beautiful than the effect presented by a mass of rough rubies when glowing in a vacuum; they shine as if they were red-hot, and the illumination effect is almost equal to that of the diamond under similar circumstances.

Masses of artificial ruby in crystals, prepared by M. C. H. Feil, behave in the vacuum like the natural ruby.

In the spectroscopic the alumina glow shows one intense and sharp red line less refrangible than the line B, and a faint continuous spectrum ending at about B. The wavelength of the red line is 6895.

The paper concludes with some notes by Prof. Maske-lyne, on the connexion between molecular phosphorescence and crystalline structure.

The crystals experimented on have been the diamond, emerald, beryl, sapphire, ruby, quartz, phenakite, tinstone, hyacinth (zircon), tourmaline, andalusite, enstatite, minerals of the augite class, apatite, topaz, chrysoberyl, peridot, garnet, and boracite. Of these, the only crystals which give out light are the diamond, ruby, emerald, sapphire, tinstone, and hyacinth. The light from emerald is crimson, and is polarised, apparently completely, in a plane perpendicular to the axis. Sapphire gives out a bluish grey and a red light polarised in a plane perpendicular to the axis. The ruby light exhibits no marked distinction in the plane of its polarisation.

Among positive crystals tinstone glows with a fine yellow light, polarised in a plane parallel to the axis of the crystal. So far the experiments accord with the quicker vibrations being those called into play, and therefore is a negative crystal the extraordinary, and in a positive crystal the ordinary, is the ray evoked. Hyacinth, however, introduces a new phenomenon, being dichroic; the colours, in three different crystals, being pale pink and lavender-blue, pale blue and deep violet, and yellow and deep violet-blue, polarised in opposite planes.

The only conclusion arrived at is that the rays, whose direction of vibration corresponds to the direction of maximum optical elasticity in the crystal, are always originated where any light is given out. As yet, however, the induction on which so remarkable a principle is suggested cannot be considered sufficiently extended to justify that principle being accepted as other than probable.

VOLUMETRIC ESTIMATION OF SULPHURIC ACID, TANNIN, &c.

By ARTHUR G. HADDOCK, A.I.C.

In the volumetric estimation of sulphuric acid by a standard solution of baric chloride I have found that by the usual method of spotting on a black glass plate, it is often extremely difficult to tell exactly when the operation is finished. It is still more so in the estimation of tannin by standard gelatine solution, especially when the extract is highly coloured.

I find that by using a small glass mirror instead of the black plate this difficulty is entirely obviated, and that the point of complete precipitation is well and sharply defined—one drop of the standard solution in excess producing a very distinct reaction. It matters not how highly

* Like the hands of a watch.

* *Annales de Chimie et de Physique*, 3rd series, vol. lvi., p. 50.

coloured the solution may be, on the mirror it appears perfectly colourless.

With the mirror it is quite easy to detect the presence of lime by means of ammoniac oxalate in one drop of the Liverpool water, even in the cold. With baric chloride the sulphates make themselves very palpably visible; indeed, I think they might almost be estimated quantitatively in the undiluted water.

Royal Institution Laboratory, Liverpool,
April 3, 1879.

RELATIONS BETWEEN TEMPERATURE AND VOLUME IN THE GENERATION OF OZONE, WITH DESCRIPTION OF A NEW FORM OF OZONATOR.

By ALBERT R. LEEDS, Ph.D.

THE methods usually employed to generate ozone, by means of the slow oxidation of phosphorus partly immersed in water, are quite unsatisfactory. Thus Miller (*"Elements of Chemistry,"* 2nd ed., Part II., p. 23) directs that a stick of clean phosphorus, moistened with a few drops of water, should be placed in a bottle of atmospheric air. In an hour or two the production of ozone attains a maximum, when, if the phosphorus be not removed, the ozone disappears, owing to its combination with the phosphorus. Instead of a bottle a large glass balloon is preferably employed, which Arenat (*"Lehr. der Anorgan. Chem.,"* p. 416) directs to be covered with a glass plate, and allowed to remain before using for twelve hours. Gorup-Besanez (*"Anorgan. Chem.,"* p. 358) recommends, in addition to the foregoing, that the bottom should be maintained for several hours at a temperature of 16° to 20°. As means of studying, or even of exhibiting to a class, the properties of ozone, everyone who has used these and similar devices will probably have found them disappointing. At times a considerable evolution of ozone occurs, at others little or none. The causes of these variations will appear on examination of the experiments detailed later.

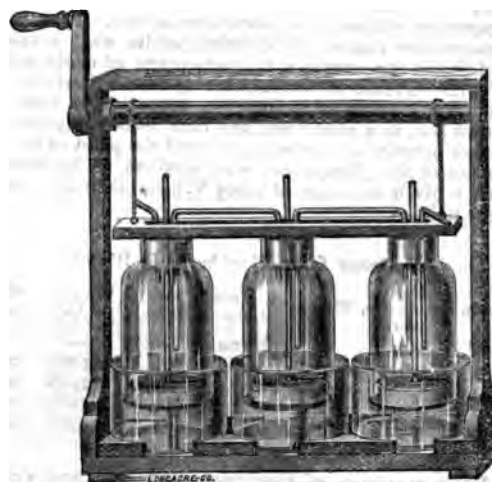
In a work published much earlier than the two last quoted, an apparatus is described and illustrated (*"Leçons de Chimie,"* A. Riche, tome i., p. 106), which was a step in the right direction. The balloon was provided with an entrance and exit tube, the latter extending nearly to the bottom, so that a slow continuous current of air might be drawn through by means of an aspirator. The ozonised air was washed in the usual manner, and the liquids to be subjected to the action of ozone were placed in a second wash-bottle. This contrivance was adopted, a large bell-jar turned upside down and covered with a glass plate being substituted for the wash-bottle when it was desired to ozonise dyed goods, flowers, and similar large objects. The apparatus was further improved by using a bell-jar in place of the balloon. The bell was set in a dish containing a number of sticks of phosphorus, the water with which they were partly covered acting as a seal. The disagreeable operation of occasionally cleansing the phosphorus by scraping was obviated by immersing them before using in a mixture of potassium bichromate solution and sulphuric acid.

These improvements, however, were of small importance compared with that effected by substituting the bichromate mixture for the water in the generator itself. The scanty fitful evolution of ozone was replaced by a copious and constant supply. The disk was now replaced by a jar capable of holding a considerable amount of the bichromate solution, and by permitting a considerable change of level in the liquid the air could either be drawn through by aspiration or forced through by pressure. Instead of placing the phosphorus on the bottom the sticks were now supported on a glass disk, which dropped into a cell connected with a paraffined iron rod sliding up

and down through the cork at top. In another generator the cell was replaced by a brass stirrup electroplated with gold; but this had to be abandoned, the solution speedily eating away the metal.

A difficulty now arose from the danger of inflammation, resulting from the great energy of oxidation. This danger seemed largely due to irregular melting down of the phosphorus, ridges being formed which protruded too far above the liquid. This unequal action also greatly diminished the amount of surface which could be safely exposed. To obviate these difficulties watch-glasses were placed in a shallow tin dish, filled with water, sufficient fragments of phosphorus placed in each, and the temperature raised until by melting the phosphorus a number of rounded cakes of uniform size had been made. Six of these could be placed upon the stage at one time, and the stage adjusted so that each cake exposed about 9 c.m. of surface area. As the convexity of the exposed portions was small, the liquid by surface action was constantly drawn on it in thin streams, and not only kept the phosphorus wet, but always clean and in condition of maximum chemical activity.

FIG. 1.



It was found, as will be seen by results stated below, that the air ozonised in one bell could have its ozone percentage notably increased, by passing through a second bell similar to the first, through a third, and so on. But as the manipulation of three, or even two, bells connected with glass tubes was troublesome, it became necessary to arrange suitable machinery. The bells were therefore cemented into heavy brass caps. Through these caps and the paraffined cork beneath, connecting tubes of heavy glass were passed, and a sliding glass rod which terminated below in a horizontal circle. The latter carried a disk of sheet lead, with a slot permitting its easy removal. The caps were then screwed fast to a board, having a suitable opening down its middle to permit of passage of tubes, &c., and this again fastened to chains by which it might be readily raised or lowered. The frame is made of such height that the jars may be easily slipped from under the bells when desirable. Thermometers are fastened into the side of each bell, their bulbs dipping just below the surface of the liquid.

It was evidently essential to convenient working of this apparatus to have a flexible connection between exit-tube and wash-bottle; but herein arose a very serious difficulty, india-rubber being destroyed. Through suggestion of Prof. Silliman, and by kindness of Mr. A. G. Day, the

* These ozonators are manufactured by S. H. Warkidge, successor to Walls and Co., Stevens Institute of Technology.

patentee, I became provided with a great variety of kerite tubing. Weighed samples of each kind were subjected to equal amounts of ozone for equal intervals, their changes and the products of decomposition noted. Without detailing these experiments, suffice it to say that the specimens which gained most in weight underwent greatest decomposition, some indeed crumbling to pieces, while others which gained none also did not change in appearance or physical characters. Of these the most satisfactory was selected, and from it as a guide Mr. Day manufactured the ozone-resisting kerite which is now used in connection with the oxonator.

The improvements effected in the ozone-generating apparatus were due, as has been said above, to a series of quantitative trials, the result of each set of experiments suggesting modifications in the next series. In the first place phosphorus was used with water alone, one bell only being employed. The ozonised air was washed, and afterwards drawn through two Peligot tubes containing a 10 per cent solution of potassium iodide. The second tube was added as a guard, but in practice is unnecessary, complete absorption occurring in the first. The liberated iodine, after acidifying with sulphuric (free from nitrous) acid, was titrated with sodium hyposulphite. In each experiment 9 litres of air were drawn over. It soon became evident that widely differing results were obtained at different temperatures, the percentage of ozone falling off with decrease of temperature to zero. Starting from this point it appeared to increase, according to some unknown law, to a maximum, and then decrease again with further increase of temperature, until the point of inflammation of the phosphorus was obtained. The volumetric ratio is given in terms of 0.005 V. p.c., taken at a convenient unit:—

I. Ozone from Phosphorus in Water,

Temp.	M. grm. per Litre, Air.	Wt. p.c.	C.c. per Litre.	Vol. Ratio.	Wt. p.c. of O.	V. p.c. of O.
20°	None	—	None	—	—	—
19.5	0.615	0.0478	0.290	5.8	0.2065	0.139
25.0	0.882	0.0682	0.413	8.3	0.2946	0.198
25.0	1.050	0.0812	0.494	9.9	0.3508	0.237
30.5	0.326	0.0252	0.153	3.1	0.1088	0.073

For convenience of comparison between these experiments and others, in which oxygen, not air, has been ozonised, columns 6 and 7 have been added—the former giving the percentage by weight of ozone in the oxygen passed over, disregarding the nitrogen; the latter, the percentage by volume. It will be seen that the maximum was attained at about 25°, when the air contained 1 m.grm. ozone in the litre, or about $\frac{1}{4}$ c.c. In the second series of experiments the water was replaced by a solution, containing to the litre of saturated solution potassium bichromate, 150 c.c. H_2SO_4 . In each experiment 8½ litres air were drawn over. The results were as follows:—

Temperature. Degrees.	Time. Minutes.	Hyposulphite. C.c.
7	—	0.40
13—14	70	4.25
20	56	6.77
21—22	—	7.10
24	60	8.35
30—31.5	60	7.13
31—32	60	7.18

From which it would appear that at a temperature of about 6° no ozone is given off, and from this point the percentage rises until the temperature attains about 24°, the percentage falling off quite evenly on both sides of the maximum. The maximum production was 1.86 m.grm. per litre air, corresponding to 0.87 c.c.

II. Ozone from Phosphorus in Bichromate. (One Bell Jar.)

Temp.	M. grm. per Litre, Air.	Wt. p.c.	C.c. per Litre.	Vol. Ratio.	Wt. p.c. of O.	V. p.c. of O.
7.0°	0.089	0.0069	0.042	0.8	0.0298	0.020
13.5	0.945	0.0730	0.441	8.8	0.3154	0.210
20.0	1.505	0.1163	0.705	14.1	0.5011	0.336
21.5	1.580	0.1221	0.741	14.8	0.5270	0.353
24.0	1.857	0.1438	0.870	17.4	0.6221	0.415
31.5	1.590	0.1229	0.745	14.9	0.5314	0.355

A third series of experiments was instituted to determine whether the air which had been ozonised in this manner could receive an increment of ozone by being again subjected to the influence of phosphorus in a second bell-jar. The liquid contained, to each litre of saturated solution of bichromate, 250 c.c. H_2SO_4 .

Temp.	Amount of Air.	Time.	Hyposulphite.
13°	2.0 litres	—	1.20 c.c.
19	2.0 "	12 minutes	2.00 "
21	2.0 "	—	2.10 "
23	3.0 "	—	3.00 "
24	3.5 "	20 minutes	4.00 "
24	2.0 "	—	2.60 "
27	2.0 "	—	2.25 "

III. Ozone from Phosphorus in Bichromate (Two Bells).

Temp.	M. grm. per Litre, Air.	Wt. p.c.	C.c. per Litre.	Vol. Ratio.	Wt. p.c. of O.	V. p.c. of O.
13°	1.134	0.0877	0.531	10.6	0.3789	0.255
19	1.890	0.1460	0.885	17.7	0.6307	0.425
21	1.985	0.1546	0.929	18.6	0.6696	0.446
23	1.890	0.1460	0.885	17.7	0.6307	0.425
24	2.155	0.1701	1.009	20.2	0.7344	0.489
24	2.457	0.1933	1.151	23.0	0.8338	0.552
27	2.127	0.1643	0.996	19.9	0.7098	0.478

It will be seen that the air ozonised in two bells likewise contains a maximum percentage at 24°, but this maximum is 25 per cent higher than the corresponding amount obtained with one bell; and in general the percentage obtained by two bells is 25 per cent greater than that with one bell at any given temperature.

Before proceeding further it appeared advisable to determine whether some other liquid might be substituted for the bichromate with advantage. Alkaline liquids seemed inapplicable, owing to the danger of formation of alkaline hyposulphites and the evolution of hydrogen phosphide. Of the acids sulphuric appeared the most suitable, although even this was attended with the disadvantage of the probable formation of some sulphurous anhydride at the same time. Mixtures of potassium permanganate and similar salts with dilute acid have not as yet been tried. The permanganate would have the disadvantage of being expensive, and even when sold as chemically pure is usually contaminated with much potassium chlorate. The results obtained with a bath containing 250 c.c. H_2SO_4 to the litre of water were as follows, 2 litres of air aspirated in each experiment, the interval varying from twelve to twenty-five minutes:—

Temperature 25°	Sodium hyposulphite 1.60 c.c.
" 26	" " 2.90 "
" 25	" " 1.06 "
" 26	" " 1.30 "
" 26	" " 1.35 "

These low results made it appear probable that the phosphorus had become oxidised. It was therefore allowed to stand for some time in the bichromate mixture, and the experiment repeated:—

Temperature 27° Sodium hyposulphite 1.70 c.c.,
Although even this amount was not so high as that previously obtained under similar circumstances, at the same

temperature, when the bichromate was employed, yet it confirmed the justness of the above supposition, and caused the use of the acid alone as a bath to be definitely abandoned. In the following table the trials are given in the order as made, with the view of showing the gradual deterioration. The abnormally high result obtained for the first 26° was due to the phosphorus having been kept for an unusual length of time in contact with the air in the bells, before aspiration was begun.

IV. *Ozone from Phosphorus in Sulphuric Acid.*
(Two Bells.)

Temp.	M.grms. per Litre Air.	Wt. p.c.	C.c. per Litre.	Vol. Ratio.	Wt. p.c. of O.	V. p.c. of O.
25°	1'512	0'1169	0'708	14'2	0'5011	0'340
26	2'741	0'2119	1'283	25'7	0'9154	0'616
25	1'002	0'0775	0'469	9'4	0'3347	0'225
26	1'229	0'0950	0'575	10'5	0'4104	0'276
26	1'275	0'0986	0'597	11'9	0'4259	0'287
27	1'607	0'1242	0'752	15'0	0'5365	0'361

Another related query was, whether so concentrated a bath was essential? To settle this point a solution was made containing, to each 1250 c.c. H₂O, 150 c.c. H₂SO₄ and 15 grms. K₂C₂O₇. The determinations were purposely made at about the maximum temperature:—

Temperature 26'5° Sodium hyposulphite 2'05 c.c.
" 26'5 " " 2'18 "

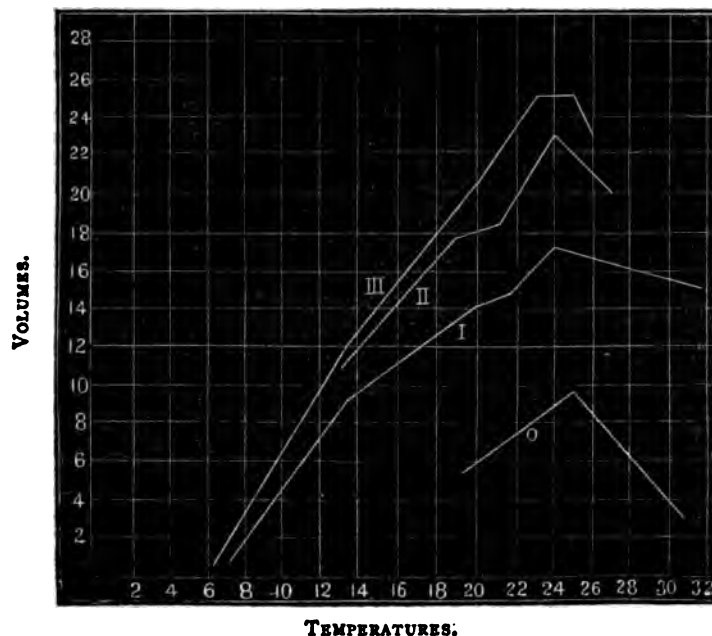
V. *Ozone from Phosphorus in Dilute Bichromate.*
(Two Bells.)

Temp.	M.grms. per Litre in Air.	Wt. p.c.	C.c. per Litre.	Vol. Ratio.	Wt. p.c. of O.	V. p.c. of O.
26'5°	1'890	0'1460	0'885	17'7	0'6307	0'425
26'5	2'060	0'1592	0'965	19'3	0'6868	0'461

These results, although inferior to those obtained with the more concentrated solution, were nevertheless so satis-

those read from the thermometers in the three bells, a reading being taken from each at the beginning, middle, and end of every experiment, and the average result stated. After the apparatus and bath had arrived at the temperature desired, the stages were raised by one motion of the crank to the height proper to expose a uniform surface of phosphorus; 8 litres of air aspirated, and then 1, 2, or more litres, the ozonised air in the latter case being titrated by solution of potassium iodide. All the figures obtained are given, although they are not so uniformly progressive as was hoped for—a result due, perhaps, in part to the difficulty of completely changing the atmosphere in three 6-litre bells by a slow current of air, without an expenditure of an amount of time which other duties would not permit.

Temperature.	Amount of Air.	Time.	Hyposulphite per Litre.
6'3°	3 litres	15 minutes	0'033 c.c.
6'3	4 "	20 "	0'033 "
10'63	2 "	15 "	0'080 "
11'1	2 "	13 "	0'100 "
11'21	2 "	15 "	0'200 "
12'22	2 "	15 "	0'225 "
12'22	2 "	13 "	0'215 "
13'33	2 "	20 "	0'680 "
13'33	2 "	10 "	0'600 "
14'99	2 "	15 "	0'400 "
14'99	2 "	15 "	0'400 "
15'88	2 "	15 "	0'420 "
16'11	1 "	10 "	0'660 "
16'11	1 "	10 "	0'680 "
17'0	2 "	15 "	0'840 "
17'5	3 "	18 "	0'610 "
20'0	2 "	18 "	1'150 "
22'78	2 "	15 "	1'520 "
22'78	2 "	10 "	1'320 "
25'0	2 "	15 "	1'420 "
26'11	2 "	13 "	1'300 "
26'11	2 "	10 "	1'200 "



TEMPERATURES.

factory and constant that this mixture was adopted, and during the subsequent work the results arrived at on increasing the number of bells to two had been so gratifying that it was thought advisable to employ three, and to combine them in an arrangement which would permit them to be easily handled, the form finally adopted being that figured in the text. The temperatures given are

An inspection of the diagram shows that in every case the maximum evolution of ozone was at or near 24° C., or about 75° F. To obtain the best results in every case, therefore, a copper tank in which the jars are placed has been added to the apparatus as figured, so that by heating or cooling the water it contains the jars and apparatus can be always maintained at the proper temperature.

VI. *Ozone from Phosphorus in Dilute Bichromate.*
(Three Bells.)

Temp.	M. grms. per Litre in Air.	Wt. p.c.	C.c. per Litre.	Vol. Ratio.	Wt. p.c. of O.	V. p.c. of O.
6.3°	0.0624	0.0048	0.029	0.6	0.0280	0.014
10.63	0.1512	0.0116	0.071	1.4	0.0501	0.034
11.1	0.1890	0.0146	0.089	1.8	0.0631	0.043
11.21	0.3780	0.0292	0.177	3.5	0.1262	0.085
12.22	0.4160	0.0321	0.195	3.9	0.1391	0.093
13.33	1.2850	0.0993	0.602	12.0	0.4290	0.289
13.33	1.1340	0.0876	0.531	10.6	0.3784	0.255
14.99	0.7560	0.0587	0.354	7.1	0.2536	0.170
15.88	0.7940	0.0618	0.372	7.4	0.2678	0.179
16.11	1.2850	0.0993	0.602	12.0	0.4290	0.289
17.0	1.5880	0.1229	0.745	14.9	0.5309	0.355
17.5	1.1530	0.0889	0.540	10.8	0.3840	0.259
20.0	2.1700	0.1677	1.018	20.4	0.7245	0.486
22.78	2.6840	0.2075	1.257	25.1	0.8964	0.603
25.0	2.6840	0.2075	1.257	25.1	0.8964	0.603
26.11	2.4570	0.1902	1.151	23.0	0.8212	0.552
26.11	2.2680	0.1778	1.062	21.2	0.7690	0.510

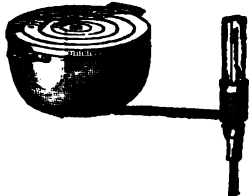
ON SOME NEW FORMS OF APPARATUS.

By M. BENJAMIN, Ph.B.

A NEW form of constant level water-bath has recently been devised, which is so simple that I will call public attention to it.

Much inconvenience has been experienced by chemists, who, having occasion to leave their laboratories for a while, on returning find that their baths have become dry, and possibly that the contents of the vessel over the bath may have been transferred to the floor by the increased heat, and the analysis is thus destroyed. The present ingenious device is so constructed as to obviate any such difficulty. Its mode of working is simple, and is as follows:—A large vessel (a common glass bottle will answer) is filled with water, and placed on a shelf or support above the bath. The water flows through a piece of rubber hose to the larger glass tube on the right of the

FIG. 1.



figure; running down this tube, it passes to the bath through the copper tube connected with the bottom of the bath. Within this glass tube is a smaller one, which is on an exact level with the top of the water-bath, through which any excess of water passes away and falls to the ground. The flow of water from the bottle can easily be regulated by means of an ordinary pinch-cock.

This water-bath is finding considerable use among students in the different laboratories, who are liable to be

FIG. 2.



called away to lectures, so that much of their time, otherwise lost, is saved by the use of this appliance.

The second engraving represents an improved form of burette clamp, which consists of two arms, one of which

is fixed, while the other is held in position by means of a joint, and is controlled by a spring, so that it acts as a lever, and by pressure it may be opened to any desired width. The end of the clamp is provided with rubber cushions, so as to hold a glass burette, or perhaps a funnel, which is placed between its arms. This clamp is made to slide on the upright of a common retort-stand.

New York, March 20, 1879.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, April 3, 1879.

Dr. WARREN DE LA RUE, President, in the Chair.

THE minutes of the Anniversary and last Ordinary Meetings were read and confirmed.

The following certificates were read for the first time:—
J. Fletcher, W. J. Bayne.

THE PRESIDENT then called on Dr. TILDEN to read a paper on "*Terpin and Terpinol*." This communication is a continuation of a previous paper (*Chem. Soc. Journ.*, June, 1878). *Crystallised Terpin*.—The liquid left after the deposition of the crystals has gone on some time contains terpinol and a nitrate, or the compound of terpinol with a nitrate, for after washing thoroughly with water and distilling in steam the yellowish oil obtained gives ammonia by the action of reducing agents, and when heated evolves nitrous fumes. Attempts to substitute sulphuric, acetic, and hydrochloric acids for nitric acid failed. The terpin hydrate, $C_{10}H_{18}O$ of Deville, seems to be nothing but terpinol. The author has obtained crystals of terpin hydrate from essence of lemon, identical in form with those obtained from American and French turpentine: he has not prepared similar crystals from the terpene of orange oil. *Terpinol*.—The author has taken the vapour density of this substance, and from it concludes that its formula is $C_{10}H_{18}O$. The alcoholic solution of terpinol dissolves 1 atom of sodium from sodium-amalgam, forming a white pasty substance, from which terpinol is regenerated by the action of water. No ether was obtained by the action of acetic acid. Hydrochloric acid gas is absorbed by terpinol, and the mixture at first turns purple, but finally is converted into a mass of colourless crystals ($C_{10}H_{18}Cl_2$), melting at 48° . From various considerations the author considers the character of terpinol to be that of an alcohol rather than that of a ketone like pinacolin. Sulphuric acid when heated with terpinol produced dehydration with partial polymerisation of the resulting hydrocarbon, which is apparently identical with terpinene. When, however, terpinol is mixed gradually with nearly unequal volume of sulphuric acid, diluted with half its bulk of water, but little heat is evolved, and no separation of the liquids takes place: on adding 3 or 4 volumes of water the whole solidifies in a few minutes into a crystallised mass of terpin, mixed with only a small quantity of liquid hydrocarbon. Oil of lemon, cajuput oil, oil of coriander and citronella apparently contain bodies either identical with terpinol or mere physical modifications of that substance. Resin spirit does not apparently contain terpinol.

Dr. ARMSTRONG thought that the reactions of terpinol pointed to a resemblance in constitution to pinacolin rather than to an alcohol, and suggested that these compounds should have a rational name expressing to some extent their constitution.

Dr. TILDEN, in reply, said that he should be reluctant to change the name until homologues were discovered, as it had been suggested by Berzelius: on the whole he adhered to his statement in the paper, that the behaviour of terpinol resembled that of a secondary or tertiary alcohol.

Mr. G. ATTWOOD then read a paper "*On a Gold Nugget from South America.*" In the State of Guayana, Venezuela, a large area of alluvial soil has lately been found to contain gold, and nuggets up to 25 ounces have been discovered within 3 feet of the surface. Numerous gold-bearing quartz veins are found in the neighbouring hills. Quite one-half of these nuggets are covered with a dark brown substance, resembling a silicate of iron. Such a nugget was treated with hydrochloric acid (its weight diminished, after treatment with HCl and NaO, from 304.7 grains to 284.33 grains). The solution contained—Silica, 0.12 gr.; ferric oxide, 8.88; lime, 0.15; magnesia, 0.08. The nugget was then treated with caustic soda, and again with HCl. The solution contained—Silica, 4.66 gr.; ferric oxide, 4.60; lime, 0.21. During this process much gold in a finely divided state became detached, and after the treatment the nugget was partly covered with a coating of finely divided gold, of a dull colour. The nugget contained 94.54 per cent of gold. The gold from the quartz veins contains 87.9 per cent gold. From these experiments the author concludes that gold nuggets gradually increase in size, owing to the accumulation of fresh particles of finely precipitated gold. Specimens of these nuggets showing the dark glazed coating were exhibited, including one weighing over 14 ounces.

Mr. W. W. FISHER then communicated a paper "*On Lead Tetrachloride.*" The existence of this compound has been for many years assumed on theoretical grounds, but direct experimental evidence has not been hitherto obtained to establish its composition. The author has therefore followed a plan similar to that already used by him to prove the existence of manganese tetrachloride. Lead dioxide dissolves in moderately strong hydrochloric acid, forming a yellow solution smelling strongly of chlorine. This yellow solution gives a precipitate of brown hydrated peroxide of lead when treated with solutions of the fixed alkalis or alkaline carbonate, &c. The addition of water causes a similar precipitation if an excess of hydrochloric acid be avoided, and the liquid carefully saturated with the dioxide. The author gives his method of analysis: he concludes that the yellow solution contains a compound of lead with chlorine, containing 1 atom of lead to 4 of chlorine. If red-lead be substituted for the lead dioxide a similar yellow solution is obtained; it can also be prepared by the action of chlorine on lead chloride suspended in dilute hydrochloric acid or solution of a chloride. If water alone be used, lead dioxide is deposited simultaneously with the formation of the yellow solution. From his experiments the author concludes that lead tetrachloride is unstable in the presence of water alone, but may exist as a double salt in the presence of other chlorides. In conclusion, the author suggests the use of chlorine or bromine in the presence of sodium acetate as a means of quantitatively determining lead by precipitation, as peroxide increases where the use of sulphuric acid is unadvisable, and gives results obtained. By thus precipitating a solution of lead acetate (in the presence of sodium acetate) as peroxide, igniting the latter to protoxide, and weighing 54.71 and 54.67 per cent Pb were obtained, theory indicating 54.67.

The PRESIDENT remarked on the importance of the suggestion as to the use of Br and Cl for precipitating lead from solution, as in many cases sulphuric acid did not completely precipitate lead.

After a short discussion, in which Drs. Wright and Tilden and Messrs. Hartley and Nelson took part, Mr. Fisher replied.

The SECRETARY then read a short note by Messrs. DALZ and SCHORLEMMER, "*On the Transformation of Aurin into Trimethyl-pararosanilin.*" This transformation was effected by the action of an aqueous solution of methylamin, at 125°, on aurin, a purple colour being formed possessing all the properties of a trimethyl-roosanilin. Intermediate compounds, soluble in alkalis, were simultaneously produced, and are at present under investigation.

The next paper was "*On the Solution of Aluminium Hydrate by Ammonia, and a Physical Isomeride of Alumina,*" by C. F. CROSS. The author has made quantitative experiments with solutions of ammonia of various strengths. His results indicate that ammonia dissolves to a certain extent the hydrated oxide at the moment of precipitation: the quantity dissolved bears no relation to the strength of the ammonia, but is considerably lessened by the presence of ammoniacal salts. The author has also examined the precipitate obtained by boiling the ammoniacal solution of the oxide: it is granular, is slowly dissolved by boiling hydrochloric acid; dried at 100° it is an opaque white powder; on igniting it undergoes no apparent change, but in the anhydrous condition it is extremely hygroscopic, absorbing 35.70 of water; its composition is Al_2O_3 .

The last communication was entitled "*Researches in Dyeing (Part II., Note on the Emission of Colouring-Matter),*" by Dr. MILLS and Mr. CAMPBELL. The experiments were made as described in Part I. (*Chem. Soc. Journ.*, 1879). The colour selected was well-crystallised Nicholson's blue. The vats contained 0.2780 grm. of pure silk ribbon, and 0.0100 grm. of blue in 400 c.c. of water; one contained in addition 1 grm. of HCl, and a third 1 grm. of NaCl. The most interesting result obtained by the authors is that the energy of combination between silk and the blue, when water or potassic chloride is used, is over-developed at first, and the excess of colouring-matter so taken up is gradually emitted by the silk during the third and fourth days; the addition of sodic chloride completely prevents this result. The authors also affirm that a real and uniform dyeing effect can always be obtained with silk and Nicholson's blue, the heat and souring used by dyers being unadvisable. The authors recommend the addition of common salt to the vat.

The PRESIDENT, in adjourning the meeting to April 17th, said that as so many candidates were awaiting election he hoped that Fellows of the Society would make every effort to be present on that date, so that a ballot might be taken.

The following papers were announced:—"On the Determination of Tartaric Acid in Lees and Inferior Argol, with some Remarks upon Filtration and Precipitation," by B. J. Grosjean. "On Conditions affecting the Equilibrium of certain Chemical Systems," by M. M. P. Muir.

CORRESPONDENCE.

ARSENICAL WALL PAPERS.

To the Editor of the Chemical News.

SIR,—I have read and heard of a good many cases of reputed poisoning from arsenical wall papers, but have never till now met with a case myself. It would seem that some people are more susceptible than others. I have slept for months in a room the paper of which contained an enormous quantity of As without experiencing any ill-effects therefrom, although I could easily rub the colour off with my finger—but to my case.

A lady in this town complained of a peculiar feeling of constriction in her nose, accompanied by a "nasty smell," nausea, and loss of appetite, and was unable to discover the cause. The symptoms had been felt since sleeping in a certain room, which she had done for two months. Somebody, I believe, suggested arsenical paper. I examined the paper and found a very large quantity of As, more than I have ever found in any paper before. H_2S gave at once a brilliant orange precipitate. Under the circumstances I could do no less than recommend that the paper should be removed, but I wanted her to try the experiment of sleeping in another room for a time to see if the symptoms disappeared, and then return again to see

if they came back, as I thought they might possibly be due to other causes. However, she would not.

I analysed the paper and found 0.35 grm. As_2O_3 per square foot, but that will scarcely represent the total amount because the paper was wetted in order to remove it, and the greater proportion of the bright green flowers washed off. There was very little copper present. I may mention that the lady's husband was not affected in the slightest degree.—I am, &c.,

A. PERCY SMITH.

Temple Observatory, Rugby
April 3, 1879.

PURE CAUSTIC POTASSA.

To the Editor of the Chemical News.

SIR,—With reference to the methods for the manufacture of potassium iodide as described by E. Schering (CHEMICAL NEWS, vol. xxxix., p. 118), he very justly ascribes the chief objection to the process in which iodine is introduced into caustic potassa and fused with carbon, to be the difficulty of obtaining caustic potassa of sufficient purity, the presence of sulphate and soda salts being most objectionable. A caustic potash practically free from these and other impurities has hitherto been unobtainable at a commercial price, and the introduction of such an article suitable for the manufacture of potassium iodide and other industries has long been a desideratum in the chemical world. This difficulty has but lately been removed by the manufacture on the large scale of caustic potash at St. Helens, by the Greenbank Alkali Works Company, of a purity which leaves nothing to be desired, a sample which recently came under my notice possessing total impurities under one per cent, of which only 0.2 was in the form of sodium hydrate. The removal of the difficulty of obtaining a caustic potash of the necessary purity being removed, the process as described above for the manufacture of potassium iodide appears to me to be the simplest and best, and having at one time been engaged in the manufacture of the article, I can therefore speak with some experience.—I am, &c.,

A. R. GARRICK, Ph.D., F.I.C.

St. Helens, April 3, 1879.

OBITUARY.

J. M. MERRICK.

WE regret to announce the death of Mr. J. M. Merrick, of Boston, U.S., who was an occasional contributor to our columns. Mr. Merrick died on the 25th day of February last, at the early age of forty-one years. In his professional life he had gained the respect of all who knew him by his uprightness of character and the ready courtesy with which he imparted his knowledge to other members of the chemical profession.

An Electric Blowpipe.—M. Jamin.—The author remarks that the electric arc which plays between two carbon conductors is a true current. If submitted to the influence of a neighbouring current, of a solenoid, or of a magnet, it experiences an action regulated by the laws of Ampère, identical with that experienced by any metallic conductor put in its place, but as its mass is exceedingly trifling its speed is considerable. The author takes advantage of this fact to submit small quantities of matter to an intense heat. By causing the arc to be driven upon lime, magnesia, or zirconia the light is directed downwards, and its intensity is increased at least three-fold.—*Comptes Rendus*.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 11, March 17, 1879.

New Experiments on Telephones without Diaphragm.—M. Ader.—The author's experiments demonstrate the truth of the opinion of M. du Moncel concerning the origin of the sounds reproduced in the telephone.

New Combinations of Hydrochloric Acid with Ammonia.—L. Troost.—The author remarks that hydrochloric acid and ammonia have hitherto been obtained only in the proportions which form sal-ammoniac, analogous to common salt. No hydrochlorate of this chloride has yet been discovered, nor an ammoniac hydrochlorate with excess of base. He has discovered a great number of curious compounds formed by dry ammonia with hydrochloric acid, hydrosulphuric acid, and other acids, both mineral and organic. He takes ammoniacal gas absolutely dry and free from every trace of compound ammonias, saturates it with pure dry hydrochloric acid, and submits the sal-ammoniac thus obtained and distilled in a close vessel to the action of a large excess of gaseous ammonia, refrigerating to different degrees. He thus obtains two well-defined compounds, characterised by their point of fusion, their crystalline structure, and their tension of dissociation. The former of these, tetra-ammoniac hydrochlorate, HCl_4NH_3 , melts at $+7^\circ$, and its crystals depolarise light powerfully. The other compound, hepta-ammoniac hydrochlorate, HCl_7NH_3 , melts at -18° .

Combinations of Hydrogen Phosphide with Cuprous Chloride and its Determination in Gaseous Mixtures.—J. Riban.—The compound $\text{Cu}_2\text{Cl}_2 \cdot 2\text{PH}_3$, chloride of cuproso-diphosphonium, is obtained by slowly passing hydrogen phosphide into a solution of cuprous chloride, refrigerated by means of ice. Hydrogen phosphide present in gaseous mixtures may be determined by agitation along with a solution of cuprous chloride in hydrochloric acid.

The Crystalline Forms of the Compounds of the Stanmethylys and of their Homologues.—M. Hiortolahl.—Not susceptible of useful abstraction.

New Method of Treating Iron and Copper Pyrites in the Dry Way.—L. Simonin.—An account of J. Hollway's experiments.

The State in which the Precious Metals are found in some of their Combinations, in Ores, Rocks, and Artificial Products.—E. Cumenge and Edmond Fuchs.—The authors find that in certain pyrites gold is found not native but in combination with antimony, and is incapable of being taken up by mercury.

Constitution of Coal.—E. Guignet.—The author on treating dry powdered coal with phenol has extracted as much as 4 per cent of a brown matter. If coal in very fine powder is treated in a cohobator with nitric acid, oxalic acid and trinitro-resorcin are distinctly recognisable among the products. Resorcin was not detected in the coal.

Alcoholic Fermentation.—P. Schützenberger and A. Destrem.—If yeast is placed in such conditions as to prevent its development and multiplication it nevertheless preserves its power of decomposing sugar, and if it acts upon sugar it de-assimilates more nitrogen than if preserved in presence of water without sugar and oxygen.

Determination of Glucose in Blood.—P. Cazeneuve.—The author criticises the method of determination followed by C. Bernard as the progress of the reduction of

the cupro-potassic liquid is often uncertain when the reaction approaches its limit. It is also reduced by other principles present in the blood. The author thinks that glycaemia should be studied anew when a more accurate analytical procedure has been devised.

Derivatives of Normal Methyl-oxyl-butyric Acid.—E. Du villier.—An account of the methyl-oxyl-butyric acid of ethyl and methyl and of methyl-oxyl-butyramid.

Nature of the Albumen of Hydrocele.—M. Bechamp.—Three distinct albumens are present different from blood albumen.

Modifications of the Physical Properties of Starch.—F. Musculus.—Amylaceous matter may exist either in the colloid or the crystalloid state. When crystalline it is soluble even in cold water.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 15, 1879.

The Chloranilins.—F. Beilstein and A. Kurbatow.—An account of the di-, tri-, and tetra-chloranilins. The authors propose to found the nomenclature of these compounds on the relative position of the chlorine atoms.

Para-di-propyl-benzol and Certain of its Derivatives.—H. Körner.—An account of sulpho-para-di-propyl-benzolic acid with its lead, barium, and calcium salts; of dinitro-para-di-propyl-benzol, and of propyl-benzoic acid.

Determination of Vapour-Densities.—V. Meyer.—Not intelligible without the accompanying engraving.

Occurrence of Furfural in Commercial Glacial Acetic Acid.—V. Meyer.—The author has found that glacial acetic acid, represented as containing from 99 to 100 per cent of real acid and apparently of unexceptional quality, assumed a splendid red colour on contact with aniline, but lost this property on distillation with chromic acid. This colouration was due to the presence of furfural of which 0.108 gm. was found per litre of acid.

Variability of the Colouring-matter of Red Wine.—J. Erdmann.—The author traces decided differences in the behaviour of genuine wine of one and the same growth according to its age.

Spectroscopic Researches on the Constitution of Solutions.—H. Burger.—In this paper the author communicates merely his methods of observation.

Malabar Gum Kino and a New Compound, Kinoin.—C. Etti.—The author added kino to twice its weight of boiling dilute hydrochloric acid (1 : 5). Kino-red separates out as a soft mass, which slowly hardens on cooling, while the kinoin remains in solution, contaminated with a little kino-red. When purified it crystallises in colourless well-formed prisms, sparingly soluble in cold water, readily at a boil, and very easily in alcohol of every concentration. The solution is not precipitated by glue and is coloured red by ferric chloride. The composition of kinoin is $C_{14}H_{12}O_6$.

Mono- and Diphenyl-arsenic Compounds.—W. La Coste and A. Michælis.—The readiness with which phosphorphenyl chloride can be obtained from phosphorus chloride and benzol induced the authors to examine if this reaction could not be generalised so as to serve for the preparation of the corresponding arsenical compounds.

Tri-phenyl-arsin and its Derivatives.—W. La Coste and A. Michælis.

Mono-tolyl-arsenic Compounds.—W. La Coste and A. Michælis.—These two papers may be regarded as continuations of the foregoing.

Schizomycetic Fermentations.—An examination of the fermentations of erythrite, glycerin, mannite, citrate of lime, malate of lime, &c., infected with cow-dung, infusion of hay, and certain varieties of purulent matter.

Thioglycolic and Thiodiglycolic Ethers.—C. Böttger.—The author has obtained these products by the

action of sulphuretted hydrogen upon glyoxylic acid in presence of silver oxide.

Mineral Waters near Buda Pest.—M. Ballo.—This memoir may possibly be of medical interest.

Studies in the Naphthalin Series.—Raphael Meldola.—The object of this communication is to throw light upon the constitution of the naphthalin derivatives. The author gives an account of dibrom-acet-naphthalide.

Sulph-etheric Acids of the Phenols.—E. Baumann.—In describing the sulpho-cresolic acids, the author remarks that the sulpho-para-cresolate of potassium is a constant ingredient in the urine of horses and probably also of other mammals.

Bromo-acetic Ethyl-ether.—F. Kessel.—A lengthy memoir, not adapted for abstraction.

Action of Dry Ammonium Sulphate upon Dry Barium-ethyl Sulphate in Presence of Baryta.—H. Köhler.—The author obtains ethyl-amin, but in small quantity only.

Decomposition of Ethyl-sulphates by Gaseous Hydrochloric Acid.—H. Köhler.—The products formed are free sulphuric acid, barium sulphate, and chlor-ethyl.

Tropæolin as an Alkalimetric Indicator.—Dr. Lunge.—The author confirms the statements of Miller concerning the use of tropæolin OO for the titration of sodium carbonate, and for determining the free acid in alumina. Not merely the carbonate but sodium sulphide (in the crystalline state) may be accurately titrated in the cold with sulphuric acid, using tropæolin as indicator. On the slightest excess of acid appearing, the yellowish colour changes at once to a magenta and then to orange, which after a few seconds disappears entirely. Tropæolin OOO undergoes the reversed play of colour, being yellow in acid solution, but turning to a magenta-red in alkalis.

Reimann's Färber Zeitung,
No. 11, 1879.

Two new green dyes have been introduced into practical use; the *Vert acide* of A. Poirrier, of Paris, and the Victoria green of the Baden Aniline Company. The price of the latter is only 16s. per kilo.

The red azo-colouring-matters are fast superseding cochineal in the flannel dye works of Saxony. They have the advantage of not being injured by washing.

No. 12, 1879.

Washing Wool with Ether and Alcohol.—The result of the experiments made at Ensival, near Verviers, is stated as follows:—70,000 kilos. wool and 16,000 kilos. yarn were extracted. The former yielded 15,800 kilos. of oil, or 22.6 per cent, and the latter 2400 kilos., or 15 per cent. The oil was sold at 39 to 40 francs per 100 kilos. The loss of ether amounted to 4.9 kilos. per 100 kilos. of wool treated.

No. 23, 1879.

It appears that milk is dyed in the cows. If fed with *Anchusa tinctoria* they yield a blueish milk, if with *Rheum palmatum* a yellowish, and if with madder or bed-straw (*Galium*) a reddish.

Justus Liebig's Annalen der Chemie,
Band 196, Heft 1.

Constitution of Phenanthren.—Gustav Schultz.—The author infers that phenanthren-quinon and consequently phenanthren are to be regarded as diortho-compounds.

On Phenanthren-quinon.—R. Anschütz and Gustav Schultz.—An account of the preparation of this compound and its behaviour with aqueous alkalis and lime, with alcoholic potassa, and alcoholic ammonia.

Researches on Hydrogen Peroxide.—E. Schöne.—The author has investigated the behaviour of hydric peroxide with the oxygen compounds of thallium. He finds that thallium paper is turned brown by the vapour of hydric peroxide in consequence of the formation of thallic oxide. Hence the browning of thallium test-papers on exposure to atmospheric air is by no means a proof of the presence of ozone.

On Aurin.—R. S. Dale and C. Schorlemmer.—The authors describe their conversion of "red corallin" or "peonin" into rosanilin. They have further examined ammonia-aurin, tetrabrom-aurin—which latter compound is very similar to tetrabrom-rosolic acid, soluble in alkalies with a fine violet colour, whilst its acidified solutions dye silk and wool a dark violet. They have further investigated the action of acetyl chloride and anhydrous acetic acid upon aurin, the compounds of aurin with acids. The compounds of aurin with acids are red like the aurin itself, whilst those with the acid sulphites of the alkali metals and with anhydrous acetic acid are colourless like leukaurin. The homologue of aurin rosolic acid likewise forms with acids well crystallised compounds, and as a powerful base. Hence it should not be considered as an acid, and might be more appropriately named rosaurin.

On Pyruvic Acid.—Dr. C. Böttger.—The author shows that the quantity of pyruvic acid obtained from glyceric acid is so trifling that it is inadmissible to make use of this manner of formation for the deduction of a constitutional formula for the latter acid. The demonstrated identity of the sulpho-lactic acid from pyruvic acid and from α -chlor-propionic acid is evidence for the ketonic character of pyruvic acid.

Communications from the Chemical Laboratory of the University of Kasan.—These consist of papers on Allyl-dipropyl-carbinol, by P. and A. Saytzeff; and on Allyl-di-ethyl-carbinol, by A. Schirokoff and A. Saytzeff.

Action of Tertiary Butyl-iodide on Isobutylene in Presence of Metallic Oxides.—J. Lermontoff.—Not susceptible of useful abstraction.

Tetra-methyl-ethylen and its Derivatives, and the Chemical Structure of Pinakon.—D. Pawlow.—Not suitable for abstraction.

La Lancette Belge.

M. Schnetzer, Professor at Lausanne, finds that in presence of borax all the colouring-matters of plants except chlorophyll are diffused, so that the plant becomes colourless; red flowers, for instance, become green.

Mr. Edison is said to have invented an ink which gives raised characters upon paper, capable of being read by the blind by touch.

MEETINGS FOR THE WEEK.

TUESDAY, 15th.—Civil Engineers, 8.
WEDNESDAY, 16th.—Meteorological, 7.
THURSDAY, 17th.—Chemical, 8. "On the Determination of Tartaric Acid in Lees and Inferior Argol, with some Remarks upon Filtration and Precipitation," by B. J. Grosjean. "On Conditions Affecting the Equilibrium of Certain Chemical Systems," by M. M. P. Muir.
Zoological, 4.

ERRATA.—P. 97, col. 1, line 6 from top, for 25 to 30 per cent read 0.25 to 0.30 per cent. Line 14 from top read 0.25 to 0.26 per cent.

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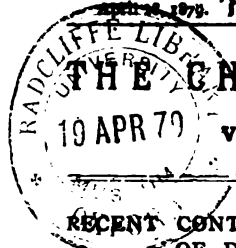
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THE CHEMICAL NEWS.

VOL. XXXIX. No. 1012.

RECENT CONTRIBUTIONS TO THE HISTORY OF DETONATING AGENTS.

By Professor ABEL, C.B., F.R.S.

AMONG the many explosive preparations which have during the last thirty years been proposed as substitutes for gunpowder, on account of greater violence and other special merits claimed for them, not one has yet competed with it successfully as a propelling agent, nor even as a safe and sufficiently reliable explosive agent for use in shells; for industrial applications and for very important military or naval uses, dependent upon the destructive effects of explosives, it has had, however, to give place, to a very important extent, and in some instances altogether, to preparations of gun-cotton and nitro-glycerin.

But there appeared little prospect that either gun-cotton or nitro-glycerin, whether used in their most simple conditions or in the forms of various preparations, would assume positions of practical importance as explosive agents of reliable, and therefore uniformly efficient, character, until the system of developing their explosive force through the agency of a detonation, instead of through the simple agency of heat, was elaborated.

Before the first step in this important advance in the application of explosive agents was made by Alfred Nobel, about twelve years ago, the very variable behaviour of such substances as gun-cotton and nitro-glycerin, when exposed to the heat necessary for their ignition under comparatively slight modifications of attendant conditions (*e.g.*, as regards the completeness and strength of confinement or the position of the source of heat with reference to the main mass of the material to be exploded) rendered them uncertain in their action, and at any rate, only applicable under circumstances which confined their usefulness within narrow limits. The employment by Nobel of an initiative detonation, produced by the ignition of small quantities of mercuric fulminate or other powerful detonating substances, strongly confined, for developing the violent explosion, or detonation, of nitro-glycerin, opened a new field for the study of explosive substances, and the first practical fruit was the successful application of plastic preparations of nitro-glycerin and of compact forms of compressed gun-cotton, with simplicity and certainty, to the production of destructive effects much more considerable than could be accomplished through the agency of much larger amounts of gunpowder, applied under the most favourable conditions. Whereas very strong confinement has been essential for the complete explosion of these substances, so long as the only known means of bringing about their explosion consisted simply of the application of fire or sufficient heat, no confinement whatever is needed for the development, with certainty, of a decidedly more violent explosive action than they are capable of exerting when thus applied, if they are detonated by submitting some small portion of the mass to the blow or concussion developed by a sharp detonation, such as is produced by the ignition of a small quantity of strongly confined mercuric fulminate.

The conditions essential to the development of detonation in masses of nitro-glycerin and gun-cotton, or preparations of them, and the relations to and behaviour towards each other of these and other explosive bodies, in their character or functions as detonating agents, has been

made the subject of study by the lecturer during the last ten years, and some of the earlier results published by him in connection with this subject also led to the pursuit of experimental inquiries of analogous character by Champion and Pellet and others.

Some of the chief results attained by Mr. Abel's experiments may be briefly summarised.

It was found that the susceptibility to detonation, as distinguished from explosion, through the agency of an initiative detonation, is not confined to gun-cotton, nitro-glycerin, and preparations containing those substances, but that it is shared, though in very different degrees, by all explosive compounds and mixtures.

It was demonstrated that the detonation of nitro-glycerin and other bodies, through the agency of an initiative detonation, is not ascribable simply to the direct operation of the heat developed by the chemical changes of the charge of detonating material, and that the remarkable property possessed by the sudden explosion of small quantities of certain bodies (the mercuric and silver fulminates) to accomplish the detonation of nitro-glycerin and gun-cotton, is accounted for satisfactorily by the mechanical force thus suddenly brought to bear upon some part of the mass operated upon. Most generally, therefore, the degree of facility with which the detonation of a substance will develop similar change in a neighbouring explosive substance, may be regarded as proportionate to the amount of force developed *within the shortest period of time* by that detonation, the latter being in fact analogous in its operation to that of a blow from a hammer or of the impact of a projectile.

Thus, explosive substances which are inferior to mercuric fulminate in the suddenness, and the consequent momentary violence, of their detonation, cannot be relied upon to effect the detonation of gun-cotton, even when used in comparatively considerable quantities. Percussion cap composition, for example, which is a mixture of fulminate with potassium chlorate, and is therefore much less rapid in its action than the pure fulminate, must be used in comparatively large quantities to accomplish the detonation of gun-cotton.

The essential difference between an explosion and what we now distinguish as a detonation lies in the comparative suddenness of the transformation of the solid or liquid explosive substance into gas and vapour.

The gradual nature of the explosion of gunpowder is illustrated, in its extreme, by burning a train of powder in the open air; the rapidity and consequent violence of the explosion is increased in proportion to the degree of confinement of the exploding charge, or to the resistance to the escape or expansion of the gases generated upon the first ignition of the confined substance. In proportion as the pressure is increased under which the progressive transformation of the explosive takes place, the rapidity with which its particles are successively subjected to the action of heat is increased.

In the case of a very much more sensitive and rapidly explosive substance than gunpowder, such as mercuric fulminate, the increase in the rapidity of its transformation, by strong confinement, is so great that the explosion assumes the character of a detonation in regard to suddenness and consequent destructive effect. A still more sensitive and rapidly explosive material (such as the silver fulminate and iodide of nitrogen) produces when exploded in open air effects akin to those of detonation; yet even with these bodies confinement operates in increasing the rapidity of the explosive to suddenness, and consequently in developing a more purely detonative action. Thus, the violence of explosion of silver fulminate is decidedly increased by confining the substance in a stout metal case, and the enclosure of iodide of nitrogen in a shell of plaster of Paris has a similar effect. With chloride of nitrogen, the suddenness of detonation, and consequently the violence of action, was found to be very greatly increased even by confining the liquid beneath a thin layer of water.

Detonation, developed in some portion of a mass, is

* Abstract of a Paper read before the Royal Institution of Great Britain, Friday, March 27, 1879.

transmitted with a velocity approaching instantaneousness throughout any quantity, and even if the material is laid out in the open air in long trains composed of small masses. The velocity with which detonation travels along trains thirty or forty feet in length, composed of distinct masses of gun-cotton and of dynamite, has been determined by means of Noble's chronoscope, and was found to range from 17,000 to 24,000 feet per second. Even when trains of these explosive agents were laid out with intervening spaces of half an inch between the individual masses composing the trains, detonation was still transmitted along the separated masses with great though diminished velocity.

The suddenness with which detonation takes place has been applied as a very simple means of breaking up shells into small fragments, and scattering these with considerable violence, with employment of very small charges of explosive agent. Thus, by filling a 16-pr. common shell completely with water, and inserting a charge of $\frac{1}{4}$ oz. of gun-cotton fitted to a detonating fuze, the shell being thoroughly closed by means of a screw-plug, the force developed by the detonation of the small charge of gun-cotton is transmitted instantaneously in all directions by the water, and the shell is thus broken up into a number of fragments averaging fourteen times the number produced by bursting a shell of the same size by means of the full amount of powder which it will contain (13 ozs.). Employing 1 oz. of powder in place of $\frac{1}{4}$ oz. of gun-cotton in the shell filled with water, the comparatively very gradual explosion of the powder charge is rendered evident by the result, the shell being broken up into less than twenty fragments by the shock produced by the first ignition of the charge, transmitted by the water. In this case the shell is broken up by the minimum amount of force necessary for the purpose, before the explosive force of the powder charge is properly developed. Extensive comparative experiments, carried on not long since by the Royal Artillery, at Okehampton, demonstrated that this simple expedient of filling common shells with water and attaching a small charge of gun-cotton with its detonator to the fuze usually employed, allowed of the application as efficient substitutes for the comparatively complicated and costly shrapnel and segment shells.

Another illustration of the sharpness of action developed by detonation as compared with explosion, consequent upon the almost instantaneous character of the metamorphosis which the explosive agent undergoes in the cases of detonation, is afforded by a method which the lecturer applied some years since for comparing the violence of action of charges of gun-cotton and of dynamite arranged in different ways. The charges (5 lbs.) to be detonated were freely suspended over the centres of plates of very soft steel of the best quality, which rested upon the flat face of a massive block or anvil of iron, having a large central circular cavity. The distance between the upper surface of the plate and the charge suspended over it was 4 feet. The sharp blow delivered upon the plate by the air suddenly projected against it by the force of the detonation when the charge was fired forced the metal down into the cavity of the anvil, producing cup-shaped indentations, the dimensions of which afforded means of comparing the violence of the detonation. A much larger charge of powder exploded in actual contact with the plate would produce no alteration of form in the metal, and the same negative result would be furnished by the explosion over the plate of a heap of loose gun-cotton of the same or greater weight than the charges detonated. The above method of experiment was devised, in the first instance, by Mr. Abel, in July, 1875, for comparing the quality of some specimens of Llandore steel proposed to be used by the Admiralty for ship-building purposes with samples of malleable iron, and it has since been employed by Mr. Adamson in carrying out a very useful series of experiments, recently communicated to the Iron and Steel Institute.

(To be continued.)

END-ON ILLUMINATION IN PRIVATE SPECTROSCOPY, AND ITS APPLICATIONS TO BOTH BLOWPIPE FLAMES AND ELECTRIC ILLUMINED GAS-VACUUM TUBES.*

By PIAZZI SMYTH,
Astronomer Royal for Scotland, and Past President of R.S.S. Arts.
(Continued from p. 146).

PART II. *Flame-Spectroscopy.*

THE various kinds of flame most frequently employed to render substances incandescent in spectroscopy, are—

- (1.) The alcohol lamp.
- (2.) The Bunsen gas burner.
- (3.) The coal-gas blowpipe driven by air.
- (4.) The coal-gas blowpipe driven by oxygen. And
- (5.) The oxy-hydrogen blowpipe.

Of these five kinds, I choose the middle one as a stand by; for while its pointed conical flame is more definite, precise, and workmanlike than the wavy loose flames of the first two, it is cheaper, more usually met with, and more frequently ready for acting, than the two latter; while it links us on to a very fair amount of heat for any flame-work, or to what we may consider a standard step in temperature throughout all these researches. The above coal-gas and air blowpipe has, moreover, the further recommendation that, dull as its blue-gray flame may appear to the eye, yet it contains several colours which separate into distinct, discontinuous bands under the spectroscopic; and they would probably be more often utilised for spectroscopic science if they could only be made decidedly brighter. But brighter it must be without any change of temperature, and also without our losing the use of the simple open flame to vapourise at pleasure adventitious and external matters in the usual flame spectroscopic manner.

Prof. Swan of St. Andrews, who first gave a large, scientific and indeed very noble account of this blowpipe flame and its spectrum, and gave it so far back as 1856, was so impressed with the importance of brightening, if possible, its faint bluish light, that he placed three small blowpipes (of his own ingenious making) one behind the other, and looked through all three flames at once. Thereby he was enabled to measure all the the leading details of four out of the really five coloured bands in this spectrum; but failed to see the lines in the first of these as to position, viz., the orange-amber band. After him several observers in London, Paris, and Manchester resorted to single or double blowpipes, but unfortunately in conjunction with pure oxygen gas; for by that emphatic addition, the temperature of their flames was so greatly raised, that they are all necessarily excluded from the present inquiry.

More recently I have obtained a copy from Paris of the late Padre Secchi's grand work entitled "*Le Soleil*" (2nd ed. 1875); and on p. 246 of vol. i.,—where that brilliant scientist has introduced, though without acknowledgment, a copy of my own Edinburgh Observatory sketch in 1872 of the whole five bands of the simple blowpipe spectrum,—he yet, in alluding to it in the text, speaks of the bands as being, in all ordinary cases witnessed by him, *three* only, and their colours "*green and blue*"—really citron, green, and blue; in which case he could only have seen the three brighter ones of the five.

In Prof. Roscoe's well-known volume of "*Lectures on Spectrum Analysis*" (1st ed. 1868), there is a coloured representation of the coal-gas spectrum with five bands, and some of them with lines; not perhaps intended to be very accurate, but most assuredly not indicating the

* Read before the Royal Scottish Society of Arts, February 20, 1879.

slightest appearance of any lines in the orange-amber band. The 3rd edition, of 1873,—which I have just referred to, contains, at p. 320, the same plate.

A more advanced style of pictorial representation is that contained in Plate III. of M. de Boisbaudran's admirable, in most of its plates unequalled, if not inimitable, work entitled "*Spectres Lumineux*" (Paris, 1874). Plate III. purports to represent the spectrum of the blue flame of coal-gas. Our five coloured bands are of course represented; but the lines, marked only in the citron, green, and blue bands, are broad and hazy; the observation having been confessedly made with a slit that was "not very narrow," and there are no lines at all on the very faint orange-amber band. Not are there any records of such things amongst the numerical measures,—though the existence of four lines is mentioned in an appendix as having just been seen when both a very strong blowpipe jet was employed, and some extraneous matters introduced into its flame, with the effect of considerably condensing it.

Such was the state of this question when, last winter, after trying many experiments with plain and simple coal-gas and air blowpipes (the coal-gas being merely taken out of the service-pipes of the house, and the air driven through its flame out of an india-rubber bag, with a pressure of 2 or 3 inches of water), it occurred to me, remarking the narrow but elongated figure of the brighter part of the flame,—viz., about $\frac{1}{2}$ inch in diameter and 2 inches long,—to try looking at it *end on*, in place of, as almost every spectroscopist had hitherto done, *transversely* to the axis of the flame; and of course quite close up to it in both cases. The result, without any other change, was almost magical; not only were five bands at once most brilliantly seen as to their colours, but the six or seven lines in the *first* of them, or the *orange-amber* band, hitherto only uncertainly made out by any one, were now the most beautiful series in the whole spectrum. Next the lines in the *citron* band showed themselves splendid for brightness, and the number of them, towards their vanishing side, much increased. In the still brighter *green* band, the first line stood forth in a manner to well deserve its quaint, Alexander Herschel title of "the green giant of carbo-hydrogen." The lines in the fainter *blue* band were yet admirably distinct, though closer than the others. And in the violet band, though no similar lines exist there, the distinction of the faint band, the strong band, and the one fine line between the two, was also well made out.

So much appeared with a very moderate prism-power on the spectroscope; but on applying a dispersion equal to 22° between A and H, or five times as much as might usually be employed, I gazed for a time in mute admiration on the scene then opened up. Such a scene of —; but, if you please, before I begin any attempt, which can hardly but prove weak and poor, to set before you some idea of what the really entrancing feature of that vision was, allow me just a minute to state why this particular spectrum is thought so very important, and is so very much sought after.

Wretched as may be the more publicly illuminating power of a grey blowpipe flame of coal-gas and simple air, it is the standard representative in spectroscopy of nearly half nature that does not, and nine-tenths of nature that does, burn. By day the spectroscopist, however far he may travel in the spectrum, from ultra red to distant lavender-grey, can always find out whereabouts he is by referring to the solar lines as seen either in the sun itself, or in his marvellous quality of light as reflected from blue sky, or a cloud, or a hill, or a house, or from snow, or from anything whatever which reflects any daylight at all. But at night, when there can be no such light to reflect,

and the last of the gloaming is gone, and there may be no moon nor planet, the young spectroscopist is either lost in darkness or bewildered on a chromatic ocean without fiducial marks of any kind. For though he turns to the light of a lamp for aid, or to any one of ten thousand gas-lights, their bright flame—bright from the incandescence of little solid particles of carbon—offers, like all other incandescent solids, only a continuous spectrum, wherein the colours blend undistinguishably one into the other from the red through the citron to the violet. Hence the poor youth at night would have been altogether at sea, when spectroscoping something totally new in heaven or on earth, had not our excellent Fellow of former years, and still only removed by the breadth of the Fifth, Prof. Swan, shown that all the substances hitherto used by man for artificially illuminating the darkness of night (such as wax, oil, tallow, turpentine, ether, alcohol, generally carbo-hydrogens)—all of them give, in the blue base of their flames, one and the same spectrum of but five coloured bands and their several innate half-dozen or so of lines; the same spectrum, in fact, as that of the flame of our favourite kind of blowpipe, though that little instrument gives it more neatly, clearly, and without the swamping effect on its best characteristics of the dense continuous spectrum derived from the more or less yellow light in the upper parts of most kinds of simple lamp-flame.

Now I had for a long time, when voyaging in the Mediterranean, actually used an alcohol lamp with a flame made purposely long and thin, in order to develop its blue base and hide its yellow top;—I had, I say, employed those carbo-hydrogen spectral lines as given by alcohol, for reference in Aurora and Zodiacal Light spectroscopy. But I gave it up at last on finding how much cleaner and freer from apparently adventitious haze the selfsame bands and their lines appeared in the blowpipe flame of coal-gas* and common air. And yet the lines did not appear even there without any haze at all! In fact, there was still so much of it hanging to them, and interfering to such an extent with the easy visibility of some of the fainter lines, that I am afraid in my haste I thought it a nuisance, an abnormal interloper, and held it as mere trash to be got rid of in any possible way. But it would not go; and now, on looking once again at the old green band and all its plaguing haze, made denser than ever by the brightness of the end-on view, but on this occasion *extra analysed, or spread out, by the tremendous dispersive power just described*—behold! all that haze, instead of becoming more hazy and diffuse than ever, was resolved neatly and perfectly into innumerable, exquisite, fine lines or linelets! These miniature lines of light, indeed, filled the whole field of view; asserted themselves to be as necessary to the spectrum of coal-gas and air as any other part of it; and had, besides, a beautifully regular progressive arrangement of their own, gradually increasing in distance asunder, as the older band lines—now looking like huge and far-between Cyclopes—decreased in theirs, as they all tended on, in the direction of increasing refrangibility, towards the violet.

Something of the same kind, and even more distinct in many of its details, has already been seen, I am aware, by many other persons, in certain spectra both of oxygen blowpipes and electric sparks; but those spectra have greatly elevated temperatures, and I am confining myself here, for the reasons already stated, to simple coal-gas and atmospheric air blowpipes only. And not so much even there, to what may be done absolutely by increasing their size immensely, as in Prof. Roscoe's blowpipe with "a blue flame 3 feet long;" but to the differential effect of looking at one and the same blowpipe flame, whether large or small, *first transversely, and then end-on*. The

* This increase of the first expected effect was owing to the hollow nature of flame, and the superficiality of its light. So that when viewed end-on, the whole of the illumination previously seen in the long side profile, was now merely condensed to a disc equal in area to a cross-section of the flame, but was further concentrated into a thin circumference of that disc.

* At a distance from gas supplies it is well to know that an equivalent to coal-gas for the spectroscope, and perfectly safe, may be produced by passing a stream of air through a vessel containing a considerable surface of benzene, or other hydrocarbon liquid in a vapourising condition at ordinary temperatures.

improvement, too, of vision for spectroscopy by the end-on method is so decided, and at the same time is obtained so simply and economically, or without any increase of expense whatever, that it would doubtless have been adopted everywhere long ago, and not left for me to advocate, had not the chemists, a large and influential body of men, and to whom we all owe much (yet said, in former times, if not still, to positively rejoice in living in the midst of flames, and smoke, and soot),—if, I repeat, the chemists had not so generally kept to their sad, dirty plan of letting their salts burn and fizz in a flame close in front of the naked slits of their spectroscopes.

Hence Mr. Rand Capron, in his "Photographed Spectra" of chemical elements, following lately the supposed orthodox method, finds himself compelled to write—"Much trouble was experienced in keeping the slit-plate clear from metallic beads and other impurities." And he suggests that in future it would be a great advantage to have the whole slit-plate gilded, and the slit-jaws formed of obsidian, platinum, or gold. Vain resource, however, if the spitting flame be still preserved close by, and quite inapplicable to any end-on use of it; for that would imply directing the full blast of the blowpipe right on the slit-plate and melting it down altogether. But my rude spectroscope having been built up at home, and primarily for operating on the distant light of the Aurora, as seen occasionally from an upper chamber at No. 15, Royal Terrace, Edinburgh, I was compelled from the first to bring such very far-off light to the slit by an anterior image-forming object-glass. And once having made experience of that excellent method of using an optical image of the thing, and not the thing itself, especially when that thing is worse to touch than a red-hot poker, have kept to it ever since for table work close by, as well as celestial objects a long way off.

The above, therefore, removing the last of the practical difficulties in the way of any one desirous of enjoying the advantages of end-on illumination in his flame spectroscopy, I close this part of the paper by referring to the Appendix for the following tables of useful data:—

Table 1. Colours of the spectrum by spectral place.

- " 2. The blowpipe flame's spectrum, seen end-on under small dispersion.
- " 3. Certain parts of the blowpipe flame's spectrum, under higher dispersion.
- " 6. Wave-number places of certain practical data.

(To be continued).

PROCEEDINGS OF SOCIETIES.

NEWCASTLE CHEMICAL SOCIETY.

(Concluded from p. 133.)

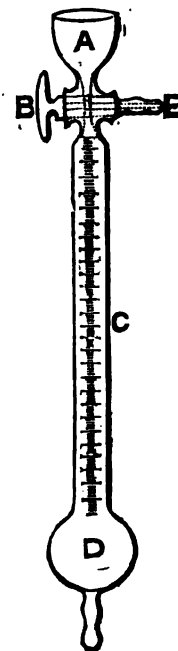
"Tennant's Nitrometer," by Messrs. BRADY and MARTIN.

This apparatus is used for testing the amount of nitrous compounds existing in sulphuric acids from the Gay Lussac and Glover towers, and also in the chamber acids. The three higher oxides are each reduced to NO, and from the volume of the latter, obtained from a given quantity of the acid, the nitrogen, nitrous acid, or nitric acid may be easily calculated.

Description.—B is a three-way stop-cock, having a passage between the tube C and the cup A, and between C and the waste pipe E. G is a tube graduated to 30 c.c. in fifths. A piece of caoutchouc tubing is fixed to the outlet of the globe D, and carried to a bottle or reservoir for mercury, from whence the graduated tube may be filled.

A measured quantity of the acid to be tested is then

introduced into the cup A, and, by lowering the mercury reservoir and opening the stop-cock, allowed to pass into the graduated tube. The stop-cock is then closed, and the column of acid vigorously shaken till its reaction on the mercury is completed.



Mr. STARKS, at the invitation of the Chairman, gave a practical description of the mode of using the nitrometers, remarking at the same time that the two instruments were identical in principle, and almost identical in detail. The instrument with the bulb was the form in use at Messrs. Tennant's works, at Hebburn, and he had used it for some time before he was aware that Dr. Lunge had contrived a similar piece of apparatus. No doubt Dr. Lunge was equally innocent of any knowledge of their doings, and the instruments were thus simultaneously invented. He admitted, however, that they were indebted to Dr. Lunge for the name—a name so obviously appropriate, that it was not desirable to invent a second.

The CHAIRMAN then described an experiment he had recently performed on the production of light, by passing a current of electricity through a slender rod of carbon enclosed in an exhausted globe, the physical results of which experiment he thought might be interesting to the members. He had obtained the current from a Siemens dynamo-electric machine of the second or 6000 candle size. On passing the current, the rod became heated to such an intense degree as to cause it to glow with great splendour. The glass became coated with a sooty deposit on its inside; and the rod of carbon, which before use had been exposed to an intense heat, to secure as much contraction as possible, became curved to the form of a bow, as if from softening.

Prof. HERSCHEL remarked that there was every appearance of the carbon having become plastic with the heat; and, unless he were right in thinking that Deville had observed the same, it was the first time carbon had been known to have been rendered plastic; but he had a strong impression that Deville had made a similar observation. The opportunity of seeing the effect, however, was very interesting.

"Halden and Thornton's Modification of the Cyanotype Process, for Copying Drawings and Tracings," by Mr. JAS. DONNELLY.

* An octavo book, with 37 remarkable plates, published by E. and F. Spoe, Charing Cross, 1877. See its page 8.

The CHAIRMAN remarked that this was but a slight modification of the Cyanotype process of Sir John Herschel, one of the earliest and simplest of the photographic processes, a process which was not adapted for artistic work, but was eminently useful for copying engineers' and architects' drawings, as it required no knowledge of chemicals or photographic art on the part of those using it; the sensitive paper being tolerably permanent, was supplied ready for use, and, after exposure, required nothing but washing in pure water to fix it.

Mr. BERKLEY said, though the process was not new, it was no less valuable, and was in constant use in the drawing office at Messrs. Palmer's works at Jarrow, and many other large establishments.

Prof. HERSCHEL confirmed what had been said by the Chairman and Mr. Berkley about the origin, the utility, and the simplicity of the process. He said it was extensively used on the Continent, and in illustration he laid upon the table a collection of very large and beautiful cyanotype prints from draughtsmen's designs which he had received from Belgium, some of the designs covering 8 to 10 square feet of sensitive paper.

"Varley's Electric Harmonium," by Prof. HERSCHEL.

It consists of a gamut of tuning-forks arranged as spring contact-breakers. A key-board is attached, and the depression of any key directs the current through an electromagnet, under the influence of which the fork applied to it vibrates, and the corresponding fork is thereupon set in similar vibration. The feeble tone of the forks was much increased when a telephone was placed in circuit, and was still further increased when the telephone was replaced by a piece of apparatus acting like a Leyden jar, and consisting of a sheet of thin gutta-percha, with perforated zinc on one side, and thin tin-foil on the other.

Mr. REYNOLDSON exhibited an automatic arrangement for the continuous supply to an evaporating-basin of a large given quantity of water; also a model of an arrangement for indicating the flow of acid through a Glover or Gay-Lussac tower. As these attracted considerable attention, and were not accompanied by a note, Mr. Reynolds was requested to bring them under the attention of the Society at some future meeting.

"Reagent Bottles with Embossed Labels." Messrs. BRADY and MARTIN exhibited microscopes by Beck and by Zeiss, and a set of reagent bottles, the labels of which were embossed in relief on the surface of the glass, and rendered more visible by being roughed by grinding, thus forming dull letters on a bright ground.

Swan's dry photographic plates were the subject of an interesting experiment, their extreme sensitiveness enabling the Chairman to obtain two successful prints from a negative; one being exposed two seconds to the light of a common bat's-wing burner at the distance of one foot, and the other exposed to the same light for only half this time.

A vote of thanks to the exhibitors and others who had contributed information to the meeting brought the proceedings to a close.

CORRESPONDENCE.

AUSTRALIAN EUCALYPTI.

To the Editor of the Chemical News.

SIR,—My old friend, Baron Ferdinand Von Mueller, Director of the Victorian Botanical Department, has very kindly sent me the special information you wished as to the comparative analysis of some of the various Eucalypti, by which you will see that the celebrated *E. globulus* is by no means the richest in the essential oil which gives the peculiar sanitary value to this class of trees, although it

is the most "taking" in its earlier stages, from the rapidity of its growth and fulness of foliage.

"*Eucalyptus Amygdalina*."

"It is this species which yields more volatile oil than any other tested, and which therefore is largely chosen for distillation; thus it is also one of the best for subduing malarian effluvia in fever regions, although it does not grow with quite the same ease and celerity as *E. globulus*. The respective hygienic value of various Eucalypts may to some extent be judged from the percentage of oil in their foliage, as stated below, and as ascertained by Mr. Bosisto at the author's instance, for the Exhibition of 1862:—

<i>E. amygdalina</i>	3'313	per cent volatile oil
<i>E. oleosa</i>	1'250	" "
<i>E. leucoxylon</i>	1'060	" "
<i>E. goniocalyx</i>	0'914	" "
<i>E. globulus</i>	0'719	" "
<i>E. obliqua</i>	0'500	" "

"The lesser quantity of oil of *E. globulus* is, however, compensated for by the vigour of its growth and the early copiousness of its foliage. The proportion of oil varies also somewhat according to locality and season. *E. rostrata*, though one of the poorest in oils, is nevertheless important for malarian regions, as it will grow well on periodically inundated places, and even in stagnant water not saline. *E. oleosa* (F. v. M.), from the desert regions of extra-tropic Australia, might be reared on barren lands of other countries for the sake of its oil. According to Mr. Osborne's experiments, Eucalyptus oil dissolves the following, among other substances, for select varnishes and other preparations:—Camphor, pine-resins, mastich, elemi, sandarac, kauri, dammar, asphalt, xanthorrhæa-resin, dragon's blood, benzoe, copal, amber, anime, shellac, caoutchouc, also wax, but not gutta-percha. These substances are arranged here in the order of their great solubility. The potash obtainable from the ashes of various Eucalypts varies from 5 to 27 per cent. One ton of the fresh foliage of *E. globulus* yields about 8½ lbs. of pearl-ash. A ton of the green wood, about 2½ lbs.; of dry wood, about 4½ lbs. For resins, tar, acetic acid, tannin, and other products of many Eucalypts see various documents and reports of the writer, issued from the Melbourne Botanic Garden."

I have had so many enquiries from different parts of the world for seeds, and as to what sort will suit various soils and climates, that I have asked Messrs. C. J. Cresswell and Co., seedsmen, of Sydney, to make a special point of securing pure seed of various kinds of trees, and from various parts of the Colony, so as to suit either tropical or cold climates (as whilst we never see frost in my garden near Sydney, yet at Kiandra the snow lies for months!); and I trust, therefore, that in a few years' time, with a little patience and experience, the right sorts to thrive, either in temperate England or torrid India, may be accurately known, and reliable plantations thus secured.

The Baron (at Melbourne) and Mr. Charles Moore, F.R.L.S., Director of our Sydney Botanical Department, will both be happy to afford any special information to parties really requiring it for practical use; but as amongst the three of us we have already answered over one hundred letters on the subject, I trust that the information therein given will be "passed on" by the recipients as much as possible, so as to save us a little, as no leading Government officials or men of business in these Colonies have any spare time worth speaking of.—I am, &c.,

R. D. ADAMS.

Sydney, N.S.W., February, 1879.

DYEING.

To the Editor of the Chemical News.

SIR,—In the otherwise accurate report of "Researches in Dyeing: Part II.," which appeared in the CHEMICAL NEWS (vol. xxxix., p. 161), Mr. Campbell and myself are repre-

sented as considering "the heat and souring used by dyers" to be "unadvisable." The statements actually made in the papers were as follows:—"The heat and souring to which dyers usually have recourse are questions quite apart from the intrinsic deposition of the pigment." . . . "We think it advisable not to maintain this blue so long as is usually done at the dissociation temperature." For HCl read KCl.—I am, &c.,

EDMUND J. MILLS.

Andersen's College, Glasgow,
April 11, 1879.

LOSS OF NITRE IN VITRIOL MANUFACTURE.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxix., p. 198, I noticed a letter from Dr. G. Lunge, in which he asserts that all fear of any appreciable loss of nitrous compounds in the Glover tower has vanished.

I have opposed Dr. Lunge on a former occasion by publishing in *Dingler's Journal* (vol. 227, p. 465, also vol. 228, p. 545), an account of some experiments on the subject of denitration, and a statement showing the loss of nitrate of soda as incurred at Messrs Gaskell, Deacon, and Co.'s works. Since that time I have continually watched the loss of nitrate of soda at these works, and applied the most refined methods of ascertaining these losses. The result of this labour has been confirmatory of all the figures I published at that time.

Chance has now presented me with the results obtained at many of the largest English alkali works. I am not at liberty to publish these. They are the results of daily tests of the exit gases extending over more than one month. The samples were all drawn by continuous aspirators and were analysed by the same method in each work. The volumes were corrected for temperature and pressure. These results and a few generally acknowledged facts enable me to make the following statements with regard to the loss of nitre in the manufacture of sulphuric acid.

I have for convenience divided the losses into two classes. The first class includes losses due to mere mechanical causes, such as the loss incurred by leakage, by the withdrawal of nitrous anhydride from the system either by imperfectly denitrated vitriol or by the exit gases. The second class I called chemical losses, because they are due to the reduction of the higher oxides of nitrogen to either nitrous oxide or nitrogen.

I. Mechanical Losses.

(a.) *Nitrous Anhydride withdrawn from the System by Means of Exit Gas.*—Taking the average amount of oxygen, leaving the Gay-Lussac column at 10 per cent, and the average amount of nitrate of soda used at 4 per 100 of sulphur passing into the chambers, the results of seven different large works show that the loss of nitre from this source is about 10.5 per cent of the nitrate used.

(b.) *Loss of Nitrous Anhydride by Imperfect Denitration.*—This loss is acknowledged by all authorities to be small. Mr. Davies (CHEMICAL NEWS, vol. xxxvii., p. 155) allows from 10 to 15 per cent of nitrate as lost by vitriol which was not at all denitrated. In Messrs. Gaskell, Deacon, and Co.'s works the loss from this source hardly ever reaches 6 per cent. From analyses of Glover-tower vitriol, published by Mr. Maclear, I calculate a loss of 2 per cent; Mr. Davies usually finds traces only in Glover-tower vitriol. If I assume 10 per cent of loss from imperfect denitration I greatly exaggerate.

(c.) *Loss from Leakage of Apparatus.*—This loss cannot be estimated directly. The loss of sulphur from all sources does not exceed 10 per cent. Most of this is lost as sulphurous acid, passing into the chimney with the exit gas, and into the air from the kilns. The amount leaking into the air from the chambers is trifling. Hence the nitrogen acids leaking into the air from the chambers

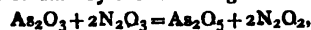
are trifling in amount also; but I will allow 5 per cent of nitrate of soda for this loss.

II. Chemical Loss.

This loss is incurred in the Glover tower, in the chambers, and, according to Mr. Davies, in the Gay-Lussac column, though not there in form of reduction to nitrous oxide. This chemical loss is exceedingly difficult to trace and to measure. As regards the chambers Mr. Davies's paper proves (if it proves anything at all) that the amount of nitre lost in this way in chambers not connected with Gay-Lussac and Glover columns is usually from 10 to 15 per cent. But allow 20 per cent for systems which are connected with such columns and you will find that:—

The exaggerated total losses from all sources amount to 45 per cent, leaving 55 per cent of the nitrate of soda used still to be accounted for. These 55 per cent represent the chemical loss from the Glover tower, or the Gay-Lussac tower, or both.

Mr. Davies thought he had explained this loss by the oxidation of the arsenious acid to arsenic acid in the Gay-Lussac column by the following reaction:—



and he gave it as his opinion that the N_2O_2 escaped into the chimney. But this explanation will not hold good, for two reasons: (1.) Because this reaction does not take place even at temperatures (220° to 230° F.) which never obtain in Gay-Lussac columns. (2.) Even if the reaction did occur to an infinitesimal amount, it does not follow that the nitric oxide is lost, since the gas usually contains sufficient oxygen (8 to 10 per cent) to re-convert it into higher oxides; and any escape of such higher oxides is already included in the mechanical loss (a).

The mere fact that As_2O_5 occurs in Gay-Lussac vitriol can be explained without having recourse to the imaginary reaction on which all Mr. Davies's calculations are based.

The loss of nitre from chemical causes cannot therefore be explained as taking place in the Gay-Lussac column, and, since only the Glover tower remains, I cannot agree to any such statement, that all fear of appreciable loss of nitre in that piece of apparatus has vanished.

It is not, in my opinion, any service to the manufacturer to put him off his guard and prevent further search by such statements.—I am, &c.,

FERDINAND HURTER.

Laboratory of Gaskell, Deacon, and Co.,
Widnes.

CHLORIDE OF CALCIUM.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxix., p. 122, there is an answer from Messrs. Gaskell, Deacon, and Co., to my notice on chloride of calcium, which you kindly published (p. 97). Allow me to add a few more words.

Very little knowledge of chemistry would be required for any person to know that chloride of calcium is a by-product in the manufacture of carbonate of soda by the Leblanc process and of bleaching-powder by the Weldon process. I wished to call the attention of manufacturers to the production of calcium chloride, because it is produced in large and increasing quantities in the manufacture of carbonate of soda by the ammonia process, which quantities are to be added to those produced by the other manufactures. Messrs. Gaskell, Deacon, and Co., advertise the sale of this product. I never doubted that it is sold by them; but I should like to know at what price these gentlemen used to sell it before the introduction of the Solvay process, which has considerably lowered its market value. It is on account of the low price and the large production that I wished to hear of new processes in which it could be used.—I am, &c.,

O. GLUGE.

Sarrebruck, April 11, 1879.

MANUFACTURE OF PICRIC ACID.

To the Editor of the Chemical News.

SIR,—I noticed in the abstract from the *Chemiker Zeitung* (No. 10, 1879), which appeared in the *CHEMICAL NEWS*, vol. xxxix., p. 150, on the above subject, that a certain J. Marzelli proposes "to add slowly the sulphacid of phenol to concentrated nitric acid." I may just mention that I know of at least one English manufacturer who used this method and manner of operating advantageously some ten years ago. So long ago as this I have also myself used the method, allowing the sulphacid to fall drop by drop through a tap-funnel into the concentrated nitric acid. This plan is an excellent one, and I am under the impression it is better and more widely known to manufacturers than J. Marzelli is aware of. I am under the impression, too, that I have seen, about a year and a half ago, more than one student in the Technical Laboratory of the Polytechnicum of this town preparing picric acid according to the above-mentioned scheme, as that of a known technical process.—I am, &c.,

WATSON SMITH, F.C.S., F.I.C.

Zürich, April 6, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 16, 1879.

Action of Anhydrous Hypochlorous Acid upon Ethylen.—E. Mulder and G. J. W. Bremer.—The authors obtained a compound represented by the formula— $C_4H_6Cl_2O_2$.

Constitution of Ultramarine.—Arthur Lehmann.—Not susceptible of useful abstraction.

Double Salts of Calcium Sulphate with other Salts and Behaviour of Gypsum in Certain Saturated Saline Solutions.—R. Fassbender.—A precipitate which the author had formerly obtained by adding solid sulphate of potassa to a saturated solution of gypsum and potassic chloride proved to be a mere mixture of the two sulphates with varying quantities of potassic chloride. The latter salt is very obstinately retained by the sulphates of calcium and potassium as is also potassic nitrate. The author confirms Struve's result that calcio-potassic sulphate is obtained by the action of potassic chloride upon gypsum.

Action of Hydrogen upon Meta-nitro-para-trichlor-acetoluyd and Meta-nitro-para-valeryl-toluyd.—T. Friederici.—Not suitable for abstraction.

New Method of Preparing Chrysaniic Acid.—T. Friederici.—The author oxidises dinitro-acetoluyd by the action of potassium pyro-chromate along with moderately dilute sulphuric acid.

On Chlor-nitro-anilins.—F. Beilstein and A. Kurbatow.—The authors examine the nitro-dichlor-anilins with para-, ortho-, and meta-position of the chlorine atoms.

On so-called Dichlor-azo-phenol.—R. Hirsch.—As regards the preparation and properties of dichlor-azo-phenol the author confirms the statements of its discoverers, Schmitt and Bennewitz. By treatment with hydrochloric acid he obtained from it hydrochlorate of dichlor-amido-phenol.

On Cinchotenicin.—O. Hesse.—The author gives this name to an isomer of cinchotenin. It is a dark brown, perfectly amorphous mass, very brittle, and yielding a yellow powder. It is readily soluble in water, alcohol, chloroform, dilute acids, ammonia, and alkaline lyes, but

is insoluble in ether. The yellowish brown solution turns the plane of polarised light slightly to the right.

Action of Cyanogen upon Amido-benzoic Acid and Anthranilic Acid in Aqueous Solution.—P. Griess.—From the reaction of cyanogen upon metamido-benzoic acid the author obtained amido-benzoic percyanide and a new compound, cyan-carbimidamido-benzoic acid. The latter of these compounds has an acid reaction, but forms salts both with bases and acids. At elevated temperatures and on treatment with acids it proves very unstable, and a number of interesting compounds are produced by its decompositions. The action of cyanogen upon anthranilic acid in aqueous solution gives rise to bicyan-amido-phenol.

Pinacons and Pinacolins.—W. Thörner and T. Zincke.—The authors give an account of the three aceton-phenon-pinacolins.

Diphenyl-methyl-acetic Acid.—W. Thörner and T. Zincke.—This compound, $C_{15}H_{14}O_2$, crystallises from dilute alcohol in white ramified leaflets, but from pure alcohol in shining transparent cubes. It melts at 173° . Several of its salts are described.

Hydrocarbon, $C_{16}H_{12}$, from Phenyl-glycol.—A. Breuer and T. Zincke.—This hydrocarbon, on oxidation with chromic acid, is converted into a quinon, $C_{16}H_{10}O_2$, which in solution is readily polarised under the influence of light. With neutral and acid alkaline sulphites it yields well-crystallised compounds. On treatment with alkali there is formed an oxy-quinon, $C_{16}H_{10}O_3$, and with ammonia an oxy-imido-compound, $C_{16}H_{11}NO_2$.

Certain Derivatives of Phenyl-acetic Acid.—O. Stöckenius.—By the reaction of ammonia upon phenyl-brom-acetic acid the author obtains phenyl-amida-etic acid along with amygdalic acid.

Dissociation of Sal-ammoniac.—C. Böttger.—The bulb of a bulb-tube is charged with sal-ammoniac, and the tube being placed nearly horizontally, a strip of red litmus paper is placed in the upper end of the tube and a strip of blue litmus in the lower. The bulb is then heated, when ammonia, being the lighter component, escapes from the upper end and turns the red paper blue, and hydrochloric acid, being the heavier, flows from the lower end and turns the blue paper red. Success depends on the inclination of the tube.

Physiologically Active Basic Constituent of the Dita Bark (Alstonia s. Echites scholaris).—Erich Harnack.—The author's daitain, $C_{22}H_{30}N_2O_4$, is a true chemical individual, and is not to be confounded with the daitain of Grupe, a mere mixture. Its physiological action upon vertebrate animals agrees with that of curare.

On Sulpho-selen-oxy-tetra-chloride.—F. Clausnizer.—The composition of this body is SO_3SeCl_4 .

Further Methods of Formation of Sulpho-selen-oxy-tetra-chloride.—F. Clausnizer.—The author has obtained this compound by the joint action of pyro-sulphuric acid and selenium tetra-chloride, of pyro-sulphuryl-chloride and selenium tetra-chloride, and of sulphuryl-hydroxyl-chloride and selen-oxy-chloride.

Action of Sulphuryl-hydroxyl-chloride upon Chlorides of Titanium, Antimony, Tin, and Silicon.—F. Clausnizer.—The author succeeded in obtaining a double compound in the case of titanium only.

Experiments on the Preparation of Sulphuryl-hydroxyl-bromide and Sulphur-oxy-tetra-bromide.—F. Clausnizer.—The results of the experiments appear to have been negative.

Constitution of Ultramarine.—R. Rickmann.—Not available for abstraction.

Contributions to the Voluminar Law and Steric Law.—H. Schröder.—From this interesting and important paper we can select only the following results, to which the author draws more special attention:—The organic elements, C, H, O, N, have in general the same

space-filling power of one stere, though there occur, e.g., in water of hydration $=H_2O_4$, certain condensations or expansions of individual elements which, as will subsequently appear, are characteristic of organic nuclei. A second very remarkable fact is the extraordinary constancy of the voluminar constitution with which certain elements and their compounds are contained in the most different groups. A third very important fact is the simplicity of the volume-molecules of all salts. Their volume-molecules are almost always monatomic, or at most di-atomic, the latter being the case only in iron spar, calcareous spar, and sodic nitrate.

Die Chemische Industrie.

No. 1, Jan., 1879.

Manufacture of Potassium Iodide.—E. Schering.—Inserted elsewhere.

Process for the Extraction of Sulphur from Alkali-waste, Gypsum, Heavy Spar, and Sulphurous Acid, and recovery of the Earths previously Combined with Sulphur in the State of Carbonates.—Max Schaffner and W. Helbig.—This process cannot be fully described without the accessory plates. It is based upon the decomposition of calcium sulphide and magnesium chloride, which latter salt has no action upon calcium carbonate. The residue from this reaction, after the expulsion of the sulphuretted hydrogen and which contains magnesia and calcium chloride, is exposed to the action of carbonic acid, when magnesium chloride is reproduced and calcium carbonate formed.

No. 2, February, 1879.

Report on the Stassfurt Industry.—Dr. B. Bernhardt.—The improvements recently introduced aim chiefly at effecting economy in the consumption of carnallite and of fuel. It is proposed to evaporate the various liquors in closed boilers under a slight pressure. Continuous evaporation is likewise suggested, the difficulty being the removal of the salts from the heated surfaces as deposited. The conversion of potassium chloride into sulphate by double decomposition with Epsom salts is advantageously effected under Dr. Borsche's patent. The mixed salts are successively lixiviated with quantities of water insufficient for complete solution. The bulk of the magnesium chloride is removed in the first liquors. Many practical men, however, still prefer to treat the potassium chloride with sulphuric acid in a Jones salt-cake furnace.

Action of Magnesium Chloride upon Steam Boilers with reference to the Magnesia Process of Purifying Feed-water.—E. Bohlig.—The author concludes from his experiments that magnesium chloride has no corrosive action as long as a slight excess of magnesium carbonate is present.

No. 3, March, 1879.

Meeting of the Association for Promoting the Interests of German Chemical Industry.—In a petition to the Reichstag concerning the use of alleged poisonous colours, the Association point out that certain dyes, such as red corallin and aurantia, denounced as poisons, have subsequently been proved perfectly innocent. They pray that no colour may be prohibited till it has been formally pronounced dangerous to public health by a commission of specially qualified chemists and physiologists after a due scientific examination. At the sessions of February 23, 24, and 25 a long debate ensued on the protective duties to be imposed on foreign chemicals, and a tariff was agreed upon to be submitted to the Reichstag.

South American Production of Iodine.—Dr. G. Langbein.—The processes in use for separating iodine from the mother-liquors of the nitre refineries may be divided into three classes. (1.) The liquors, without previous concentration, are mixed with a quantity of sodium sulphite corresponding to the iodine present, the iodine separated out is filtered through linen bags, washed,

pressed, and sublimed. (2.) The mother-liquors are treated with sodium sulphite or bisulphite till the precipitated iodine is converted into hydriodic acid, which is then precipitated as cuprous iodide by a solution of copper sulphate and sodium sulphite. (3.) The iodine in the mother-liquors is concentrated by fractionated evaporation and crystallisation, and the iodine, mixed with a proportionate quantity of sodium sulphite or bisulphite, is recovered by distillation from the acidified liquid.

Efficacy of Compounds Employed as "Antichlore."—Dr. G. Lunge.—The author has submitted the *modus operandi* of some of these agents to a critical examination. The hyposulphite of soda (now thiosulphate) was said by Fordos and Gélis to undergo a direct conversion into sulphate. Its reaction with chlorine, as well as that of the sulphites, gives rise to free acid, which, if not at once washed out, attacks the iron of the machinery, forming injurious salts which render the paper brittle, yellow, or spotty. There is no sufficient proof that the use of hyposulphite leads to the deposition of sulphur and the formation of acids by its gradual oxidation if the half-stuff has been properly washed. The efficacy of hyposulphite is much less than has been assumed. Ammonia as an antichlore appears to act only at an elevated temperature with the emission of an unbearable odour.

Chemiker Zeitung.

No. 12, 1879.

Adulteration of Wax.—Dr. Max.—Bee's-wax is often sophisticated to the extent of 33 to 50 per cent with ceresin, a mixture of refined earth-wax and caranaba-wax. Pure bee's-wax has a higher specific gravity (0.94 to 0.97) than these ingredients or than the mixtures, which range from 0.876 to 0.937. Prof. v. Wagner recommends the sample to be placed in dilute alcohol of sp. gr. 0.945, and rejected as impure unless it floats.—*Dingler Polyt. Journal.*

Detection of Starch in Milk.—Dr. Vulpinus coagulates the sample with a few drops of acetic acid, heats to a boil, filters, and adds to the clear whey an aqueous solution of iodine, which at once produces a blue cloud if starch be present; 5 milligrams were thus distinctly recognised in 5 c.c. of milk.—*Pharm. Zeitung.*

Chloride of magnesium is recommended as a fluid for filling wet gas-meters.

No. 13, March 27, 1879.

Distillation of Coal-tar.—The author gives an account of the distinctions between the products obtained by the dry distillation of wood, peat, lignite, and coal, and shows that according to the recent experiments of Letny, Liebermann, and others, wood-tar and petroleum-tar, &c., if passed through ignited tubes, yield the same products as coal-tar, which he considers may be of great importance for the production of the aniline and alizarin colours. He then gives an account of the distillation of coal-tar with especial reference to anthracen.

At Dortmund, on March 22, Dr. Schridde delivered a lecture on "Chemistry as a Trade, an Art, and a Science."

In Switzerland the factory inspectors receive a yearly salary of 6000 francs, with an addition of 7 francs daily for their expenses when travelling on official business and 5 francs more if detained over night.

Moniteur Scientifique, Quasneville.

February, 1879.

This issue is almost entirely taken up with a translation of Sir B. C. Brodie's well-known "Chemical Calculus," a continuation of the discussion between MM. Berthelot and Pasteur, a notice of foreign chemical researches consisting entirely of extracts from the *Berichte*

der Deutschen Chem. Gesell., an account of the recent great discovery by Mr. Norman Lockyer, and a very lengthy notice of a work on applied physiology by M. L. Fignier.

Detection of Tarry Matters in Ammonia.—The well-known nitric acid test is recommended, which, in presence of aniline, toluylidn, &c., gives a gooseberry-red colour.

Ethyl-green and Malachite Green.—Another Swiss firm, M.M. Gerber and Uhlmann, of Bâle, announce that their green colour is prepared by the reaction of the hydride of benzoin upon ethyl- or methyl-aniline, and that it does not fall within the limits of the patent for "malachite green."

Archives Néerlandaises des Sciences.
Tome xliii., 4me Livraison.

Researches on Quinamin.—A. C. Oudemans.—The author first gives the analysis of the crude mixture of alkaloids known as quinetum, which he finds composed of:—

Cinchonin	37.0
Quinin	6.1
Cinchonidin	22.9
Quinamin	4.5
Amorphous alkaloids	21.1
Carbonate of soda	2.9
Water	2.7

From this mixture he separates quinamin, $C_{19}H_{24}N_2O_2$, the properties, reactions, and salts of which he describes in detail.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 11, March 13, 1879.

This issue contains no original chemical matter.

MISCELLANEOUS.

Royal Institution of Great Britain.—The following are the arrangements for the Lectures after Easter:—

Ernst Pauer.—Three Lectures on Schubert, Mendelssohn, and Schumann (with Musical Illustrations); on Tuesdays, April 22 to May 6.

Professor Dewar, M.A., F.R.S.—Fives Lectures on "Dissociation;" on Thursdays, April 24 to May 29.

H. H. Statham.—Four Lectures on the Leading Styles of Architecture Historically and Æsthetically Considered; on Saturdays, April 26 to May 17.

Professor Karl Hillebrand.—Six Lectures on the Intellectual Movement of Germany from the Middle of the Last to the Middle of the Present Century; on Tuesday, May 13; Mondays, May 19, 26, June 2; Tuesday, June 10; and Thursday, June 12.

John Robert Seeley, M.A., Professor of Modern History, Cambridge.—Four Lectures on Tuesdays, May 20, 27, June 3; and on Thursday, June 5.

Professor Henry Morley.—Three Lectures on Swift; on Saturdays, May 24 to June 7.

The following are the probable arrangements for the Friday evening Meetings after Easter, to which Members and their friends only are admitted.

Friday, April 25.—Francis Galton, M.A., F.R.S., M.R.I. Generic Images.

Friday, May 2.—Professor John G. M'Kendrick, M.D. The Physiological Action of Anæsthetics.

Friday, May 9.—Sir John Lubbock, Bart., M.P., F.R.S., M.R.I. The Habits of Ants.

Friday, May 16.—Professor A. Cornu. Etude Optique de l'Elasticité. (In French.)

Friday, May 23.—W. H. Preece, M.R.I. Multiple Telegraph, or Duplex and Quadruplex Telegraphy.

Friday, May 30.—Grant Allen. The Colour-sense in Insects; its Development and Reaction.

Friday, June 6.—Professor Dewar, M.A., F.R.S.

Friday, June 13.—Frederick J. Bramwell, F.R.S. The "Thunderer" Gun Explosion.

Another Agricultural College.—A correspondent of *The Live Stock Journal* writes as follows:—"With considerable satisfaction I learn that the starting of another Agricultural College is already under contemplation, for it has long been evident that the Cirencester institution has neither attained the position which its promoters had in view, nor conferred on the agriculture of the country the benefits which were reasonably expected from it. I hail therefore with pleasure the intimation that the four leading professors at Cirencester, viz., Professors Church, Tanner, Fream, and Sheldon, having been requested by influential persons to consider the idea, are already reducing to form the scheme of a second Agricultural College, for which, I have no doubt, they will receive all the help and patronage that are necessary to bring their project into successful operation. There is, indeed, room for several agricultural colleges, and no doubt the Cirencester one would have had a better tale to tell if two or three rival establishments had existed to spur it on. I regard the starting of a second Agricultural College in England as an event that is calculated to do a much-needed service to the parent institution, and I venture to predict that each of the two colleges will do better than the single one has hitherto done, provided the Council will seriously take in hand those changes which are evidently so much needed at Cirencester."

NOTES AND QUERIES.

Honey.—Can any of your readers give me references to any late (within ten years) investigations of the chemical composition of honey?—E. H. S.

Insurance of Tar Distilleries.—Can any of your readers give the name of any Fire Insurance Office of good standing which effects insurances on risks as hazardous as tar distilleries?—W. A. R.

Testing of Coal-Tar Products.—(Reply to T. C. W.)—The liquid products—naphtha, benzole, phenols, &c.—are tested by fractional distillation, observing the proportions by volume passing over between certain temperatures, and in the case of phenol observing also the proportion crystallising, and taking the melting-point of this. These test-proportions, the temperatures at which they shall pass over, and in the case of phenol the melting-point, are usually fixed by the trade contract. I know no book containing an account of the methods, which are simple cases of fractional distillation. Information on the testing of anthracen and the other solid products, as also of the aniline colours, &c., will be found usefully given in Bolley and Kopp's "Manuel Pratique d'Essais et de Recherches Chimiques Appliquées aux Arts et à l'Industrie." Paris: F. Savy, 24, Rue Hautefeuille.—WATSON SMITH, F.C.S., F.I.C.

MEETINGS FOR THE WEEK.

MONDAY, 21st.—Medical, 8.30.
Society of Arts, 8. "Recent Advances in Telegraphy," by W. H. Preece. (Cantor Lectures.)

TUESDAY, 22nd.—Civil Engineers, 8.
Royal Institution, 3. "Schubert," by Mr. Ernst Pauer.

WEDNESDAY, 23rd.—Society of Arts, 8. "English Fresh-water Fisheries," by J. Willis-Bund.

THURSDAY, 24th.—Royal, 8.30.
Royal Institution, 3. "Dissociation," by Prof. Dewar.

FRIDAY, 25th.—Royal Society Club, 6.30.
Royal Institution, 9. "Generic Images," by Mr. Francis Galton.

FRIDAY, 25th.—Quekett, 8.
Royal Institution, 3. "Architecture," by Mr. H. H. Statham.

PHYSICAL, 3.—"On Selective Reflection," by Capt. Abney. "On some Phenomena Connected with Magneto-electric Induction," by C. Boys. "Notes from the Physical Laboratory of University College, Bristol," by Dr. S. P. Thompson.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

MARCH, 1879.

The following are the returns of the Society of Medical Officers of Health:—

	Appearance in 2 foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	Sulphuric Acid Hydride.	Hardness on Clark's Scale.		
		Saline.	Organic.								Before Boiling.	After Boiling.	
		Grs.	Grs.								Degs.	Degs.	
<i>Thames Water Companies.</i>													
Grand Junction	Clear	0'000	0'008	0'165	0'062	21'20	8'020	0'720	1'152	2'360	13'0	4'20	
West Middlesex	Clear	0'000	0'007	0'120	0'064	21'00	8'170	0'576	1'080	2'160	15'4	3'70	
Southwark and Vauxhall	Clear	0'000	0'008	0'135	0'075	21'00	7'840	0'721	1'152	1'400	14'4	3'30	
Chelsea	Clear	0'000	0'008	0'126	0'064	19'80	7'280	0'720	1'152	1'740	14'3	3'30	
Lambeth	Clear	0'000	0'006	0'135	0'079	21'50	8'140	0'721	1'152	1'700	14'8	4'20	
<i>Other Companies.</i>													
Kent	Clear	0'000	0'002	0'360	0'004	29'20	10'920	1'081	1'800	2'730	17'6	5'10	
New River	Clear	0'000	0'004	0'150	0'037	19'40	7'400	0'620	1'152	1'670	14'3	3'30	
East London	Clear	0'000	0'008	0'126	0'064	23'80	8'840	0'940	1'200	2'100	16'5	4'20	

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

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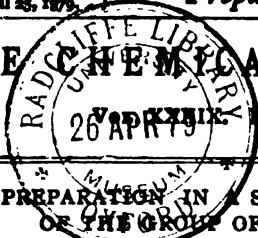
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THE CHEMICAL NEWS.



Vol. XXIX. No. 1013.

THE PREPARATION IN A STATE OF PURITY
OF THE GROUP OF METALS
KNOWN AS THE PLATINUM SERIES,
AND NOTES UPON
THE MANUFACTURE OF IRIDIO-PLATINUM.*

By GEORGE MATTHEY.

IN this paper it is not my intention, nor should I be able, to refer generally to the results of work in the various branches of platinum metallurgy carried out by my firm, who, as is well known, have been associated with the development of this special field of industry from its earliest infancy; but I shall confine myself simply to that section of it upon which my personal attention has of late years been specifically concentrated in order to meet and comply with the requisition of the Bureau Internationale des Poids et Mesures, the Section Française de la Commission Internationale du Mètre, and of l'Association Géodésique Internationale (all of them important scientific committees, formed with the object of arriving at an accurate and definite solution of the long agitated question of standard weights and measures), and also at the demand of the French Minister of War, for an alloy the best adapted for the manufacture of the international metre and kilogram standard, and the geodesique rule; and in my endeavour to solve this difficult problem I have had the great advantage of being able to consult those distinguished men, MM. Henri Sainte-Claire Deville and Henri Debray, of Paris, and have also had the benefit of the excellent and valued advice of M. Stas, the celebrated Belgian chemist, to all of whom the scientific world owe so much, and to whom I desire to offer my warmest thanks.

In a paper of this kind it would be superfluous for me to enter into any of the already published details concerning the existence and collection of what is known as platinum-dust or mineral. It is sufficient for me to observe that the six metals (of which platinum is the chief) usually found more or less in association in their native state, present characteristics of interest beyond their metallurgical utility, which are, perhaps, worth alluding to *en passant*. It is, for instance, a curious fact that the group should consist of three light and three heavy metals, each division being of approximately the same specific gravity—the heavier being (in round figures) just double the density of the lighter series.

Thus we find osmium, iridium, platinum forming the first division, of the respective specific gravities of 22.43, 22.39, 21.46; whilst ruthenium, rhodium, and palladium are represented by the figures 11.40, 11.36, 11, the average densities of the heavy and light divisions thus being respectively 22.43 and 11.25.

But a more interesting and important classification is what I may designate as a first and second class series, from the more important view of their relative properties of stability. Thus platinum, palladium, and rhodium form the first or higher class, not being volatilisable in a state of oxide; iridium, osmium, and ruthenium forming the second or lower class, their oxides being more or less readily volatilised.

The oxide of iridium is effected at 700° to 800° C., and entirely decomposed at 1000°, whilst osmic and hyporuthenic acids are volatilised at the low degree of 100°, the latter exploding at 108°. The chlorides of these metals can be sublimed at different temperatures (as also the protochloride of platinum).

I now propose to give a short description of the methods I have employed for preparing the pure platinum and iridium necessary for the manufacture of the alloy, which I call "iridio-platinum," and it is upon the distinguishing characteristics above-mentioned that my method of separation is chiefly founded.

Platinum.

The preparation of this metal to a state of purity is an operation of extreme delicacy. I commence by taking ordinary commercial platinum; I melt this with six times its weight of lead of ascertained purity, and, after granulation, dissolve slowly in nitric acid diluted in the proportion of 1 volume to 8 of distilled water. The more readily to ensure dissolution, it is well to place the granulated alloy in porcelain baskets such as are used in the manufacture of chlorine gas for holding the oxide of manganese. When the first charge of acid is sufficiently saturated, a fresh quantity should be added until no more action is apparent; at this stage the greater part of the lead will have been dissolved out together with a portion of any copper, iron, palladium, or rhodium that may have been present. These metals are subsequently extracted from the mother-liquors, the nitrate of lead by crystallisation, and the remaining metals by well-known methods.

The metallic residue now obtained will be found in the state of an amorphous black powder (a form most suitable for further treatment), consisting of platinum, lead, and small proportions of the other metals originally present—the iridium existing as a brilliant crystalline substance insoluble in nitric acid. After digesting this compound in weak aqua regia, an immediate dissolution takes place of the platinum and lead, leaving the iridium still impure, but effecting a complete separation of the platinum.

To the chloride of platinum and lead after evaporation is added sufficient sulphuric acid to effect the precipitation of the whole of the lead as a sulphate, and the chloride of platinum after dissolution in distilled water is treated with an excess of chloride of ammonium and sodium, the excess being necessary in order that the precipitated yellow double salt may remain in a saturated solution of the precipitant. The whole is then heated to about 80°, and allowed to stand for some days; the ammonio-chloride of platinum will settle down as a firm deposit at the bottom of the vessel, whilst if any rhodium, as is generally the case, is present, the surface liquor will be coloured a rose tint, occasioned by a combination of the salts of the two metals.

The precipitate must be repeatedly washed with a saturated solution of chloride of ammonium and subsequently with distilled water charged with pure hydrochloric acid. This is necessary for its purification. The small quantity of the double salt which will be taken up and held in solution is of course recovered afterwards. Rhodium may still exist in the washed precipitate, which must therefore not be reduced to the metallic state until its separation is completed, and this is best effected by mixing with the dried compound, salts of chloro-platinate and chloro-rhodate of ammonia, bisulphate of potash with a small proportion of bisulphate of ammonia, and subjecting to a gradual heat brought by degrees up to a dull red in a platinum capsule, over which is placed an inverted glass funnel. The platinum is thus slowly reduced to a black spongy porous condition free from water, nitrogen, sulphate of ammonia, and hydrochloric acid, the rhodium remaining in a soluble state as bisulphate of rhodium and potash, which can be dissolved out completely by digesting in boiling distilled water; a small quantity of platinum will have been taken up in a state of sulphate, but is regained by heating the residue (obtained on evaporation) to redness, at which heat it is reduced to the metallic condition, the rhodium salt remaining undecomposed.

By the method above described the platinum is freed not only from rhodium, but from all other metals with which it may have been contaminated, and is brought to

* A Paper read before the Royal Society.

a state of absolute purity, of the density 21.46, the highest degree obtainable.

Iridium.

In the preparation of this metal when intended to be used for the manufacture of iridio-platinum alloy, I have arrived at freeing it to the utmost possible extent from all its associate metals, except platinum, disregarding the presence of the latter; the proportion of which, once determined, would only form matter of calculation in the final operation of mixing my alloy.

In practice, the purest iridium which can be obtained from its ordinary solution (deprived of osmium by long boiling in aqua regia and precipitated by chloride of ammonium) will almost invariably contain traces of platinum, rhodium, ruthenium, and iron.

I fuse such iridium in a fine state of division with ten times its weight of lead, keeping it in a molten state for some hours, dissolve out the lead with nitric acid, subject the residue to a prolonged digestion in aqua regia, and obtain a crystalline mass composed of iridium, rhodium, ruthenium, and iron, in a condition suitable for my further treatment. By fusion at a high temperature with an admixture of bisulphate of potash, the rhodium is almost entirely removed, any remaining trace being taken up together with the iron in a later operation. The iridium so far prepared is melted with ten times its weight of dry caustic potash, and three times its weight of nitre, in a gold pan or crucible; the process being prolonged for a considerable time to effect the complete transformation of the material into iridiate and ruthenate of potash, and the oxidation of the iron; when cold, the mixture is treated with cold distilled water. The iridiate of potash of a blue tinge will remain as a deposit almost insoluble in water, more especially if slightly alkaline, and also the oxide of iron.

This precipitate must be well washed with water charged with a little potash and hypochlorite of soda until the washings are no longer coloured, and then several times with distilled water.

The blue powder is then mixed with water strongly charged with hypochlorite of soda, and allowed to remain for a time cold, then warmed in a distilling vessel, and finally brought up to boiling-point until the distillate no longer colours red, weak alcohol acidulated with hydrochloric acid.

The residue is again heated with nitre and potash water charged with hypochlorite of soda and chlorine, until the last trace of ruthenium has disappeared.

Further, to carry out the purification, the blue powder (oxide of iridium) is re-dissolved in aqua regia, evaporated to dryness, re-dissolved in water, and filtered.

The dark-coloured solution thus obtained is slowly poured into a concentrated solution of soda and mixed with hypochlorite of soda, and should remain as a clear solution without any perceptible precipitate, and subjected in a distilling apparatus to a stream of chlorine gas, should not show a trace of ruthenium when hydrochloric acid and alcohol are introduced into the receiver. In this operation the chlorine precipitates the greater part of the iridium in a state of blue oxide, which after being collected, washed, and dried, is placed in a porcelain or glass tube, and subjected to the combined action of oxide of carbon and carbonic acid obtained by means of a mixture of oxalic with sulphuric acid gently heated.

The oxide of iridium is reduced by the action of the gas leaving the oxide of iron intact, the mass is then heated to redness with bisulphate of potash (which will take up the iron and any remaining trace of rhodium), and after subjecting it to many washings with distilled water, the residue is washed with chlorine water to remove any trace of gold, and finally with hydrofluoric acid, in order to take out any silica which might have been accidentally introduced with the alkalis employed or have come off the vessels used.

The iridium, after calcination at a strong heat in a

charcoal crucible, is melted into an ingot, and after being broken up and boiled in hydrochloric acid, to remove any possible trace of iron adhering to it through the abrasion in breaking up, should possess if perfectly pure a density of 22.39; but, as iridium prepared even with the utmost care will still contain minute though almost inappreciable traces of oxygen, ruthenium, rhodium, and possibly iron, the highest density I have yet attained is 22.38.

Alloy of Iridio-Platinum.

This compound metal possesses physical properties of great value, forming a beautiful example of the effect of a careful combination of the opposite characteristics of its component parts. Thus, the extreme softness and expansiveness of pure platinum and the brittleness and excessive hardness of pure iridium, produce, by combination in judicious proportions, a perfect and homogeneous alloy, possessing the necessary mean of these properties to render it suitable for many important purposes, amongst others that of the special object to be attained to meet the requirements for an unalterable standard metal, for which it is peculiarly adapted.

In the manufacture of the prototype metres and the geodesique rules (each 4 metres in length) ordered from my firm by the Comité Internationale des Poids et Mesures, the Association Geodesique Internationale, and the French Minister of War, I proceeded in the following manner with the platinum and iridium prepared as described above.

Operating upon a charge of 450 ounces of platinum and 55 ounces of iridium, I commenced by melting these metals together and casting into an ingot of suitable shape, which I then cut into small pieces with hydraulic machinery. After re-melting and retaining in a molten condition under a powerful blast of oxygen and common gas for a considerable time, I re-cast and forged at an intense white heat under a steam hammer, the highly polished surfaces of which were cleaned and polished after each series of blows—when sufficiently reduced it was passed through bright polished steel rollers, cut into narrow strips, and again slowly melted in a properly shaped mould, in which it was allowed to cool. I thus obtained a mass of suitable shape for forging, perfectly solid, homogeneous, free from fissures or air-holes, and with a bright and clean surface at bottom and sides as at top. At the first forging a bar was obtained 35 centims. long, 7.5 wide, 2.5 thick, which weighed—

In air		15.105 grms.
In water at 70° F.		14.405 "
Showing a density at zero of	21.522	

A third of the bar was cut off and the larger portion again forged to a length of 95 centims., width 2.5, thickness 2.0, which weighed—

In air		10.814 grms.
In water at 60° F.		10.315 "
Showing a density at zero of	21.648	

This was then passed through highly polished rolls until of a length of 4080 centims., 21 millims. in width, and 5 millims. thick, to which a perfectly rectangular form was subsequently given by drawing it through a series of plates, and thus prepared the rule was in a condition to receive the beautiful polish of which this alloy is susceptible.

After passing it through each hole the metal was annealed by means of a jet of gas and oxygen to a heat just below melting-point, and each time throughout after forging, rolling, and drawing was exposed to the action of melted borax, and boiled in concentrated hydrochloric acid to remove any possible trace of adherent iron or other impurity.

A piece cut from the end and presented to the French Academy of Sciences gave the following results:—

Weight in air		116.898 grms.
" water		111.469 "
Showing a density of		21.516

thus proving that the necessary processes of annealing at a high temperature had caused it sensibly to resume its original density.

The analysis gave—

	I.	II.
Platinum	89'40	89'42
Iridium	10'16	10'22
Rhodium	0'18	0'16
Ruthenium	0'10	0'10
Iron	0'06	0'06
	00'00	00'06

From which is deduced :—

	Proportion. Density at zero. Volume.		
Iridio-platinum, at 10 p.c.	99'33	21'575	4'603
Iridium, in excess.. ..	0'23	22'380	0'010
Rhodium	0'18	12'000	0'015
Ruthenium	0'10	12'261	0'008
Iron	0'06	7'700	0'008
	99'90		4'644

Density at zero, calculated after No. 1 analysis, 21.510

"	"	"	"	No. 2	"	21'515
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thus coinciding perfectly with the practical results obtained.

Messrs. Leon Brunner Brothers, of Paris, who had submitted this material of the geodesique rule to a great number of mechanical experiments, communicated the result of their observations to M. H. Sainte-Claire Deville, thus:—

" Paris, 27 Août, 1878.

"MONSIEUR,

« La division de la règle géodésique, que nous faisons pour l'Association Géodésique Internationale, est terminée depuis quelques jours.

« Nous avons pensé que vous ne seriez pas mécontent d'apprendre que cette opération à parfaitement réussi, et que c'est au métal que nous attribuons la facilité avec laquelle nous avons pu l'exécuter.

« La platine irridie de M. Matthey est incontestablement supérieure au platine ordinaire, pour la confection de règles divisées. Il est exempt de ces pailles qu'on rencontre toujours dans ce dernier, et se laisse polir au charbon. On peut, sans danger, enlever les rébarbes des traits et les conserver très beaux. Le platine ordinaire ne peut-être poli qu'au papier à émeré, et l'on est toujours exposé à gâter la division quand on procède à l'ébarbage. C'est là un inconvénient très-grave.

"Nous ne pouvons que vous remercier, Monsieur, d'avoir mis à notre disposition un métal qui modifie singulièrement les difficultés qu'on rencontre dans la fabrication d'une règle géodésique, et nous vous prions de recevoir l'assurance de nos sentiments les plus distingués."

"BRUNNER FRERES.

In the year 1876 the suggestion was made to supersede the rectangular form by a tubular one, and I was requested to produce one of the following dimensions:—Length, 1002 centims.; exterior diameter, 37 millims.; interior diameter, 35 millims.; with rounded ends, one having an extension of small tube 4 millims. exterior diameter, 2 millims. interior diameter, 40 millims. long, which I did by the system of tube-making with automatic joints adopted by me with excellent results for the last twenty years, employing for the purpose an alloy prepared as above described. These proved to be so satisfactory that I have since made others, both round and square, of various dimensions, as lately shown at the Paris Exhibition.

Indio-platinum alloy has now been proved to possess the following among many advantages for standard rules and weights :—

It is almost indestructible, has extreme rigidity, especially in the tube form, and a most beautifully polished surface can be obtained upon it; its coefficient of elasticity is very great, whilst for standard weights its high density

is a valuable quality, and for these I should indeed recommend an alloy of not less than 20 per cent of iridium. I lately made at the request of M. H. Sainte-Claire Deville a cylinder 40 millims. by 40 millims. of such an alloy, which showed by analysis the following proportions:—

Platinum	..	..	..	..	80°6600
Iridium	..	..	..	..	19°0786
Rhodium	..	..	..	..	0°1220
Ruthenium	..	..	..	..	0°0460
Iron	..	..	..	..	0°0080

100'0046

and gave the density of 21.614.

With such a high density its coefficient of elasticity is 22·200000, one of the highest known, whilst its malleability and ductility are almost without limit.

The volume of the kilogram thus prepared is only 46266 c.c., it displaces 2267 c.c. less than the kilogram of the archives of France, and on this account, as on many others, is of course preferable.

The results I have arrived at in preparing alloys of higher grades, viz., 25, 30, 40, and 50 per cent of iridium, are as follows:—

The alloy of 20 per cent iridium is, as I have stated already, malleable and ductile.

25 per cent can only with great difficulty and waste be worked into sheet and wire when heated at low temperature. 30 per cent and 40 per cent with great difficulty only at a temperature little less than melting-point, being brittle when cold, but with a grain of great beauty and fineness.

50 per cent I have as yet failed to work up into forms other than castings beyond what I can effect by pressure when in a semi-fused condition.

The general results of my work on this alloy would lead me, therefore, to make the following recommendations.

For the manufacture of standard rules to use an alloy of not less than 85 per cent platinum and 15 per cent iridium, adopting the tubular form.

For the standard weights to use an alloy of not less than 80 per cent platinum and 20 per cent iridium, adopting the form now generally made.

Finally, following the expression of the great French chemist, M. Dumas, I hope by these labours "d'avoir enrichi l'outillage scientifique d'un alliage doué des propriétés précieuses."

RECENT CONTRIBUTIONS TO THE HISTORY OF DETONATING AGENTS.*

By Professor ABEL, C.B., F.R.S.

(Continued from p. 166.)

It has been stated that detonation can be transmitted from one mass of gun-cotton or dynamite to another through intervening air-spaces. The extent to which such spaces can be introduced without checking detonation is obviously regulated by the size of the masses of explosive detonated; but the distance of air-space through which the detonation of a moderate quantity of the explosive agent will communicate to similar masses are very limited, a space of 2 inches being sufficient to prevent the detonation produced by a mass of 8 ozs. of gun-cotton, freely exposed, from communicating to contiguous ones. If the dispersion of the force is prevented in part, and direction is given to the gases violently projected from the centre of detonation, the power of transmitting detonation to separated masses of explosive is increased to a remarkable degree. This is readily accomplished through the agency of tubes, the charge first detonated being just inserted

* Abstract of a Paper read before the Royal Institution of Great Britain, Friday, March 21, 1879.

into one extremity, while that to which the detonation is to be transmitted is inserted into the other; or separate charges may be placed at different distances inside a long tube, with long intervening spaces, the initiative charge being inserted at one end. A few illustrations of the results thus obtained may be given. The detonation of a 1-oz. disk of gun-cotton in the open air will not transmit detonation with *certainty* to other disks placed at a greater distance than half an inch from it; but if it be just inserted into one of an iron tube 2 feet long and 1.25 inch in diameter, a similar disk, or even a plug of loose gun-cotton, inserted into the other extremity of the tube, will invariably be detonated. With employment of 2 ozs. of gun-cotton, in a tube of the same material, thickness, and diameter, detonation was transmitted to a distance of 5 feet. In tubes of the same kind, of very considerable length, 2-oz. disks of gun-cotton, placed at intervals of 2 feet, were detonated through the initiative detonation of one such disk inserted into one extremity of the tube. In other experiments a long tube of this kind was fitted with branch pipes, 2 feet long, at those parts where the intermediate disks were placed, and charges of gun-cotton were placed at the extremities of these pipes. By the initiative detonation of 1 oz. of gun-cotton all the charges were detonated, the effect on the air being that of one single explosion. The results obtained with equal quantities of gun-cotton varied with the diameter, strength, and nature of the material of the tubes used. Dynamite and mercuric fulminate, applied to their own detonation, furnished results quite analogous to those obtained with gun-cotton; but in applying fulminate to the detonation of gun-cotton through the agency of tubes, some singular and instructive results were obtained, for an account of which the lecturer referred to his Memoir on this subject.

Silver fulminate was employed for the purpose of instituting more precise experiments than could be made in operating on a larger scale, with gun-cotton, on the influence of the material composing the tubes, of the condition of their inner surfaces, and other variable circumstances, upon the transmission of detonation. Half a grain of silver fulminate, freely exposed and ignited by a heated body, will transmit detonation to some of the compound placed at a distance of 3 inches from it, but does not do so with certainty through a distance of 4 inches. But when the quantity of the fulminate is just inserted into one end of a stout glass tube, 0.5 inch in diameter and 3 feet long, its detonation is invariably induced by that of a similar quantity of the fulminate placed just inside the other extremity of the tube; this result is uncertain when the length of the tubes of the same thickness and diameter exceeds 3 feet 3 inches. Glass tubes were found to transmit the detonation of silver fulminate much more rapidly than tubes of several other materials of the same diameter and thickness of substance. Thus, with the employment of double the quantity of fulminate required to transmit the detonation with certainty through a glass tube of the kind described, 3 feet in length, it was only possible to obtain a similar result through a pewter tube 31.5 inches long, a brass tube 23.7 inches long, an india-rubber tube 15.8 inches long, and a paper tube 11.8 inches long. The difference in the results obtained was not ascribable to a difference in the escape of force on the instant of detonation, in consequence of the fracture of the tube, nor to the expenditure of force in work done upon the tube at the seat of detonation, since the glass tubes were always destroyed by the first explosion to a much greater distance along their length than any of the others, and the brass tubes, which were in no way injured at the seat of explosion, did not transmit detonation to so great a distance as the pewter tubes, which were always deeply indented. The transmission of detonation appeared, also, not to be favoured by the sonorosity or the pitch of the tube employed, as the sonorous brass tube was not found to favour the transmission to the same extent as the pewter tube. Moreover, the transmission of detonation by the glass tubes was not found to be at all affected by coating these

tubes with several layers of paper, or by encasing them in tightly-fitting india-rubber tubes. These differences appeared on further investigation not to be ascribable, to any important extent, if at all, to the difference in the nature of the material composing the tubes, but to be simply, or at any rate almost entirely, due to differences in the condition of the inner surfaces of the tubes. Thus, brass tubes, the inner surfaces of which were highly polished, and paper tubes, when coated inside with highly glazed paper, transmitted the detonation of the silver fulminate to about the same distance as the glass tubes; on the other hand, when the inner surfaces of the latter were slightly roughened by coating them with a film of fine powder, such as French chalk, they no longer transmitted detonation to anything like the distance which they did when the inner surfaces were in the normally smooth condition. Other very slight obstacles to the unimpeded passage of the gas-wave through the tubes were found greatly to reduce the facility with which detonation could be transmitted by means of tubes. Thus, when a diaphragm of thin bibulous paper was inserted into the glass tube about half-way between the two extremities, detonation was not transmitted, even with the employment of about six times the quantity of fulminate that gave the result with certainty under ordinary conditions; and, similarly, the transmission of detonation by increased charges of mercuric fulminate and of gun-cotton was prevented by the introduction into the tubes of light tufts of carded cotton-wool just sufficient in quantity to shut out the light in looking through the tubes.

Among several other interesting results furnished by an examination into the conditions governing and results attending the transmission of detonation by tubes, a remarkable want of reciprocity was found to exist between mercuric fulminate and gun-cotton. The latter substance is more susceptible to the detonative power of mercuric fulminate than of any other substance, as will presently be further shown. The quality of fulminate required to detonate gun-cotton is regulated by the degree to which the sharpness of its own detonation is increased by the amount of resistance to rupture offered by the envelope in which the fulminate is confined. From 20 to 30 grains are required if the detonative agent is confined in a thin case of wood or in several wrappings of paper; but as small a quantity as 2 grains of the fulminate suffices to effect the detonation of compressed gun-cotton, provided the fulminate be confined in a case of stout metal (sheet tin), and be closely surrounded by being tightly imbedded in the mass of gun-cotton. If there be no close contact between the two, the quantity of fulminate must be very considerably increased to ensure the detonation of the gun-cotton; and, in attempting to transmit detonation from mercuric fulminate to gun-cotton by means of tubes, it was found necessary to employ comparatively very large quantities of fulminate in order to accomplish this, even through short lengths of tubes. But when the quantity of fulminate used reaches certain limits, the detonation may be transmitted from it to gun-cotton through very long lengths of tube. In applying gun-cotton, on the other hand, to accomplish the detonation of mercuric fulminate, it was found that this result could be attained, and through considerable lengths of tube (7 feet and upwards), by means of very much smaller quantities of gun-cotton than is needed of fulminate to induce the detonation of gun-cotton through the corresponding distances.

This want of reciprocity between two detonating agents corresponds to one even more remarkable, which was observed by the lecturer in his earlier investigations on this subject. In the first place it was found that the detonation of $\frac{1}{2}$ oz. of gun-cotton (the smallest quantity that can be thus applied) induced the simultaneous detonation of nitro-glycerin, enclosed in a vessel of sheet-tin, and placed at a distance of 1 inch from the gun-cotton; while with $\frac{1}{2}$ oz. of the latter the same effect was produced with an intervening space of 3 inches between the two substances. But on attempting to apply nitro-glycerin to

of gun-cotton, the quantity of the former, detonated in close contact with compressed was gradually increased in the first instance subsequently even to 2 ozs., without accommodation of the latter, which was simply a fine state of division, in all instances but one, number of experiments.

developed by the detonation of nitro-glycerin by careful comparison of the relative destruction of corresponding quantities, to be decidedly that of the fulminate, of which from 2 to 5 is for developing the detonation of gun-cotton, in close contact with them. The non-susceptibility of cotton to detonation by nitro-glycerin is need scarcely be said, not ascribable to any mechanical force suddenly applied when the is detonated.

lower possessed by different very highly explosives, of inducing the detonation of such bodies and nitro-glycerin is not solely ascribable to a of mechanical force very suddenly developed, not only by the singular inertness of gun-influence of nitro-glycerin as a detonating also by a comparison of the behaviour of other substances with that of the mercuric fulminate, d to the detonation of gun-cotton. Thus, the of silver fulminate is very decidedly sharper the mercury compound, yet it is in no way be latter in its power as an initiative detonant; indeed, a somewhat larger amount of it be required than of the mercury salt to induce of gun-cotton with certainty. Again, the chloride of nitrogen are far more susceptible of action than the silver fulminate; yet while be latter, confined in a stout metal envelope, stonate gun-cotton, 50 grains of chloride of fined by water, appeared to be the minimum i which the detonation of gun-cotton could be d with certainty, while no success attended sent of confined iodide of nitrogen in quantity up to 100 grains.

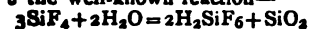
compatibility of these results with the general based upon numerous and greatly varied experiments the facility with which the detonation of and nitro-glycerin, and bodies of a similar explosives, is induced by an initiative detonation to the mechanical force aided by the ed by the latter, led the lecturer to the conclusion of synchronism or similarity in character or quality ions developed by the detonation of particular operates in favouring the detonation of one nce by the initiative detonation of a small another, while in the absence of such synchronism more powerful detonation, or the application ster force, would be needed to effect the detonation material operated upon. This view has siderable support from results since obtained erimenters, especially by M.M. Champion and the subject is one which still needs further excitation.

(To be continued.)

NEW VOLUMETRIC METHOD OF DETERMINING FLUORINE.

By SAMUEL L. PENFIELD,
Assistant in the Sheffield Laboratory.

is the well-known reaction—



f a volumetric determination of fluorine, be quantity of hydro-fluosilicic acid formed weight of a fluoride by means of a standard B.]

It is impossible to titrate the hydro-fluosilicic acid directly, because as soon as an alkaline reaction is reached the silico-fluoride is decomposed and an acid reaction is indicated, which change goes on slowly. But when barium chloride and an equal volume of alcohol are added to the solution, barium silico-fluoride is precipitated from the solution, and an equivalent amount of hydrochloric acid is liberated, which can be titrated; by this means, using litmus as an indicator, I was enabled to get some very satisfactory results, but the turbidity caused by the barium silico-fluoride interfered with the change of colour of the litmus, and cochineal I was not able to use at all.

On further experimenting I found that potassium chloride possessed some advantages over barium chloride. On adding potassium chloride and an equal volume of alcohol, potassium silico-fluoride is precipitated from the solution, and an equivalent of hydrochloric acid is liberated, but the potassium silico-fluoride is a very transparent precipitate, and does not interfere with the change of colour of the indicators. It is necessary that alcohol shall make up at least one-half the volume of the liquid to be titrated, so as to precipitate the potassium silico-fluoride as completely as possible.

The apparatus needed is very simple, and consists of a gasometer of ten or more litres capacity, a few flasks of about 150 c.c. capacity, a few large plain U-tubes 18 centimetres long and 2½ centimetres diameter, made without any narrowing in the bend, and a heavy iron plate supported properly so that it may be heated with a lamp.

The fluoride is weighed out accurately into one of the flasks; unless it is a silicate, 10 grms. of powdered and ignited quartz are added, and two or three pieces of quartz about the size of kidney beans. These last facilitate mixing up the powder when the flask is shaken. The contents of the flask are then drenched with from 30 to 40 c.c. of sulphuric acid which has been previously heated and allowed to cool. The flask is tightly closed with a doubly perforated cork; from the gasometer dry air is passed into the flask by means of a glass tube which reaches nearly to the bottom. The silicon fluoride mixed with air passes from the decomposing flask first through a small U-tube, made from ordinary glass tubing, 5 millimetres in diameter, and kept cool by being placed in a beaker of cold water, then into one of the large U-tubes intended for decomposing the silicon fluoride and absorbing the hydro-fluosilicic acid. The U-tube contains a solution of potassium chloride mixed with an equal volume of alcohol; the escaping gas is made to bubble through this, and to ensure complete decomposition a second smaller U-tube is attached to the first: the first tube absorbs nearly all the acid, the second contains only traces. The decomposing flask is supported on the iron plate; by its side is placed a second flask containing sulphuric acid and a thermometer supported so that its bulb dips into the acid; the lamp heating the plate is placed midway between the flasks, and the heat is regulated so that the temperature of the acid remains between 150° and 160° C.

The decomposition is continued two hours in ordinary cases, and during that time a continuous current of air is forced through the apparatus, amounting to from 5 to 6 litres for the two hours, while the contents of the decomposing flask are frequently agitated by shaking.

After the decomposition and aspiration are completed the contents of the U-tubes are titrated. For this purpose they may be transferred to a beaker, the tubes being rinsed with alcohol and water, or better, the acid may be titrated directly in the U-tubes. In order that the alcohol may make up one-half the volume of the liquid after the titration is completed, add a few c.c. of alcohol before titrating, or where 15 or more c.c. are to be added use a standard alkali one-half of whose volume is alcohol. A the separated silicic acid sticks to the sides of the U-tube it is necessary to have prepared a glass rod bent a little at one end to scrape off and break up this silica,

into one extremity, while that to which the detonation is to be transmitted is inserted into the other; or separate charges may be placed at different distances inside a long tube, with long intervening spaces, the initiative charge being inserted at one end. A few illustrations of the results thus obtained may be given. The detonation of a 1-oz. disk of gun-cotton in the open air will not transmit detonation with *certainty* to other disks placed at a greater distance than half an inch from it; but if it be just inserted into one of an iron tube 2 feet long and 1.25 inch in diameter, a similar disk, or even a plug of loose gun-cotton, inserted into the other extremity of the tube, will invariably be detonated. With employment of 2 ozs. of gun-cotton, in a tube of the same material, thickness, and diameter, detonation was transmitted to a distance of 5 feet. In tubes of the same kind, of very considerable length, 2-oz. disks of gun-cotton, placed at intervals of 2 feet, were detonated through the initiative detonation of one such disk inserted into one extremity of the tube. In other experiments a long tube of this kind was fitted with branch pipes, 2 feet long, at those parts where the intermediate disks were placed, and charges of gun-cotton were placed at the extremities of these pipes. By the initiative detonation of 1 oz. of gun-cotton all the charges were detonated, the effect on the air being that of one single explosion. The results obtained with equal quantities of gun-cotton varied with the diameter, strength, and nature of the material of the tubes used. Dynamite and mercuric fulminate, applied to their own detonation, furnished results quite analogous to those obtained with gun-cotton; but in applying fulminate to the detonation of gun-cotton through the agency of tubes, some singular and instructive results were obtained, for an account of which the lecturer referred to his Memoir on this subject.

Silver fulminate was employed for the purpose of instituting more precise experiments than could be made in operating on a larger scale, with gun-cotton, on the influence of the material composing the tubes, of the condition of their inner surfaces, and other variable circumstances, upon the transmission of detonation. Half a grain of silver fulminate, freely exposed and ignited by a heated body, will transmit detonation to some of the compound placed at a distance of 3 inches from it, but does not do so with certainty through a distance of 4 inches. But when the quantity of the fulminate is just inserted into one end of a stout glass tube, 0.5 inch in diameter and 3 feet long, its detonation is invariably induced by that of a similar quantity of the fulminate placed just inside the other extremity of the tube; this result is uncertain when the length of the tubes of the same thickness and diameter exceeds 3 feet 3 inches. Glass tubes were found to transmit the detonation of silver fulminate much more rapidly than tubes of several other materials of the same diameter and thickness of substance. Thus, with the employment of double the quantity of fulminate required to transmit the detonation with certainty through a glass tube of the kind described, 3 feet in length, it was only possible to obtain a similar result through a pewter tube 31.5 inches long, a brass tube 23.7 inches long, an india-rubber tube 15.8 inches long, and a paper tube 11.8 inches long. The difference in the results obtained was not ascribable to a difference in the escape of force on the instant of detonation, in consequence of the fracture of the tube, nor to the expenditure of force in work done upon the tube at the seat of detonation, since the glass tubes were always destroyed by the first explosion to a much greater distance along their length than any of the others, and the brass tubes, which were in no way injured at the seat of explosion, did not transmit detonation to so great a distance as the pewter tubes, which were always deeply indented. The transmission of detonation appeared, also, not to be favoured by the sonorosity or the pitch of the tube employed, as the sonorous brass tube was not found to favour the transmission to the same extent as the pewter tube. Moreover, the transmission of detonation by the glass tubes was not found to be at all affected by coating these

tubes with several layers of paper, or by encasing them in tightly-fitting india-rubber tubes. These differences appeared on further investigation not to be ascribable, to any important extent, if at all, to the difference in the nature of the material composing the tubes, but to be simply, or at any rate almost entirely, due to differences in the condition of the inner surfaces of the tubes. Thus, brass tubes, the inner surfaces of which were highly polished, and paper tubes, when coated inside with highly glazed paper, transmitted the detonation of the silver fulminate to about the same distance as the glass tubes; on the other hand, when the inner surfaces of the latter were slightly roughened by coating them with a film of fine powder, such as French chalk, they no longer transmitted detonation to anything like the distance which they did when the inner surfaces were in the normally smooth condition. Other very slight obstacles to the unimpeded passage of the gas-wave through the tubes were found greatly to reduce the facility with which detonation could be transmitted by means of tubes. Thus, when a diaphragm of thin bibulous paper was inserted into the glass tube about half-way between the two extremities, detonation was not transmitted, even with the employment of about six times the quantity of fulminate that gave the result with certainty under ordinary conditions; and, similarly, the transmission of detonation by increased charges of mercuric fulminate and of gun-cotton was prevented by the introduction into the tubes of light tufts of carded cotton-wool just sufficient in quantity to shut out the light in looking through the tubes.

Among several other interesting results furnished by an examination into the conditions governing and results attending the transmission of detonation by tubes, a remarkable want of reciprocity was found to exist between mercuric fulminate and gun-cotton. The latter substance is more susceptible to the detonative power of mercuric fulminate than of any other substance, as will presently be further shown. The quality of fulminate required to detonate gun-cotton is regulated by the degree to which the sharpness of its own detonation is increased by the amount of resistance to rupture offered by the envelope in which the fulminate is confined. From 20 to 30 grains are required if the detonative agent is confined in a thin case of wood or in several wrappings of paper; but as small a quantity as 2 grains of the fulminate suffices to effect the detonation of compressed, gun-cotton, provided the fulminate be confined in a case of stout metal (sheet tin), and be closely surrounded by being tightly imbedded in the mass of gun-cotton. If there be no close contact between the two, the quantity of fulminate must be very considerably increased to ensure the detonation of the gun-cotton; and, in attempting to transmit detonation from mercuric fulminate to gun-cotton by means of tubes, it was found necessary to employ comparatively very large quantities of fulminate in order to accomplish this, even through short lengths of tubes. But when the quantity of fulminate used reaches certain limits, the detonation may be transmitted from it to gun-cotton through very long lengths of tube. In applying gun-cotton, on the other hand, to accomplish the detonation of mercuric fulminate, it was found that this result could be attained, and through considerable lengths of tube (7 feet and upwards), by means of very much smaller quantities of gun-cotton than is needed of fulminate to induce the detonation of gun-cotton through the corresponding distances.

This want of reciprocity between two detonating agents corresponds to one even more remarkable, which was observed by the lecturer in his earlier investigations on this subject. In the first place it was found that the detonation of $\frac{1}{2}$ oz. of gun-cotton (the smallest quantity that can be thus applied) induced the simultaneous detonation of nitro-glycerin, enclosed in a vessel of sheet-tin, and placed at a distance of 1 inch from the gun-cotton; while with $\frac{1}{2}$ oz. of the latter the same effect was produced with an intervening space of 3 inches between the two substances. But on attempting to apply nitro-glycerin to

the detonation of gun-cotton, the quantity of the former, which was detonated in *close contact* with compressed gun-cotton, was gradually increased in the first instance to $\frac{1}{2}$ oz., and subsequently even to 2 ozs., without accomplishing the detonation of the latter, which was simply dispersed in a fine state of division, in all instances but one, in a large number of experiments.

The force developed by the detonation of nitro-glycerin was found, by careful comparison of the relative destructive effects of corresponding quantities, to be decidedly greater than that of the fulminate, of which from 2 to 5 grains suffice for developing the detonation of gun-cotton, when it is in close contact with them. The non-susceptibility of gun-cotton to detonation by nitro-glycerin is therefore, it need scarcely be said, not ascribable to any deficiency in mechanical force suddenly applied when the nitro-glycerin is detonated.

That the power possessed by different very highly explosive substances, of inducing the detonation of such bodies as gun-cotton and nitro-glycerin is not solely ascribable to the operation of mechanical force very suddenly developed, is indicated not only by the singular inertness of gun-cotton to the influence of nitro-glycerin as a detonating agent, but also by a comparison of the behaviour of other detonating substances with that of the mercuric fulminate, when applied to the detonation of gun-cotton. Thus, the detonation of silver fulminate is very decidedly sharper than that of the mercury compound, yet it is in no way superior to the latter in its power as an initiative detonating agent; indeed, a somewhat larger amount of it appeared to be required than of the mercury salt to induce detonation of gun-cotton with certainty. Again, the iodide and chloride of nitrogen are far more susceptible of sudden detonation than the silver fulminate; yet while 5 grains of the latter, confined in a stout metal envelope, suffice to detonate gun-cotton, 50 grains of chloride of nitrogen confined by water, appeared to be the minimum amount with which the detonation of gun-cotton could be accomplished with certainty, while no success attended the employment of confined iodide of nitrogen in quantities ranging up to 100 grains.

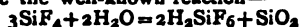
The incompatibility of these results with the general conclusion, based upon numerous and greatly varied experiments, that the facility with which the detonation of gun-cotton and nitro-glycerin, and bodies of a similar character as explosives, is induced by an initiative detonation, is proportionate to the mechanical force aided by the heat developed by the latter, led the lecturer to the conclusion that a synchronism or similarity in character or quality of the vibrations developed by the detonation of particular substances, operates in favouring the detonation of one such substance by the initiative detonation of a small quantity of another, while in the absence of such synchronism, a much more powerful detonation, or the application of much greater force, would be needed to effect the detonation of the material operated upon. This view has received considerable support from results since obtained by other experimenters, especially by MM. Champion and Pellet; but the subject is one which still needs further experimental elucidation.

(To be continued.)

ON A NEW VOLUMETRIC METHOD OF DETERMINING FLUORINE.

By SAMUEL L. PENFIELD,
Assistant in the Sheffield Laboratory.

I HAVE made the well-known reaction—



as the basis of a volumetric determination of fluorine, estimating the quantity of hydro-fluosilicic acid formed from a given weight of a fluoride by means of a standard alkali solution,]

It is impossible to titrate the hydro-fluosilicic acid directly, because as soon as an alkaline reaction is reached the silico-fluoride is decomposed and an acid reaction is indicated, which change goes on slowly. But when barium chloride and an equal volume of alcohol are added to the solution, barium silico-fluoride is precipitated from the solution, and an equivalent amount of hydrochloric acid is liberated, which can be titrated; by this means, using litmus as an indicator, I was enabled to get some very satisfactory results, but the turbidity caused by the barium silico-fluoride interfered with the change of colour of the litmus, and cochineal I was not able to use at all.

On further experimenting I found that potassium chloride possessed some advantages over barium chloride. On adding potassium chloride and an equal volume of alcohol, potassium silico-fluoride is precipitated from the solution, and an equivalent of hydrochloric acid is liberated, but the potassium silico-fluoride is a very transparent precipitate, and does not interfere with the change of colour of the indicators. It is necessary that alcohol shall make up at least one-half the volume of the liquid to be titrated, so as to precipitate the potassium silico-fluoride as completely as possible.

The apparatus needed is very simple, and consists of a gasometer of ten or more litres capacity, a few flasks of about 150 c.c. capacity, a few large plain U-tubes 18 centimetres long and $2\frac{1}{2}$ centimetres diameter, made without any narrowing in the bend, and a heavy iron plate supported properly so that it may be heated with a lamp.

The fluoride is weighed out accurately into one of the flasks; unless it is a silicate, 10 grms. of powdered and ignited quartz are added, and two or three pieces of quartz about the size of kidney beans. These last facilitate mixing up the powder when the flask is shaken. The contents of the flask are then drenched with from 30 to 40 c.c. of sulphuric acid which has been previously heated and allowed to cool. The flask is tightly closed with a doubly perforated cork; from the gasometer dry air is passed into the flask by means of a glass tube which reaches nearly to the bottom. The silicon fluoride mixed with air passes from the decomposing flask first through a small U-tube, made from ordinary glass tubing, 5 millimetres in diameter, and kept cool by being placed in a beaker of cold water, then into one of the large U-tubes intended for decomposing the silicon fluoride and absorbing the hydro-fluosilicic acid. The U-tube contains a solution of potassium chloride mixed with an equal volume of alcohol; the escaping gas is made to bubble through this, and to ensure complete decomposition a second smaller U-tube is attached to the first: the first tube absorbs nearly all the acid, the second contains only traces. The decomposing flask is supported on the iron plate; by its side is placed a second flask containing sulphuric acid and a thermometer supported so that its bulb dips into the acid; the lamp heating the plate is placed midway between the flasks, and the heat is regulated so that the temperature of the acid remains between 150° and 160° C.

The decomposition is continued two hours in ordinary cases, and during that time a continuous current of air is forced through the apparatus, amounting to from 5 to 6 litres for the two hours, while the contents of the decomposing flask are frequently agitated by shaking.

After the decomposition and aspiration are completed the contents of the U-tubes are titrated. For this purpose they may be transferred to a beaker, the tubes being rinsed with alcohol and water, or better, the acid may be titrated directly in the U-tubes. In order that the alcohol may make up one-half the volume of the liquid after the titration is completed, add a few c.c. of alcohol before titrating, or where 15 or more c.c. are to be added use a standard alkali one-half of whose volume is alcohol. A standard alkali one-half of whose volume is alcohol. A separated silicic acid sticks to the sides of the U-tube it is necessary to have prepared a glass rod bent a little at one end to scrape off and break up this silica,

I have found by experiment that a simple dry U-tube between the decomposing flask and absorbing tube is sufficient to condense any sulphuric acid that may go over from the heated acid.

When fluorine is to be determined in a mineral containing chlorine, as in the case of an apatite, substitute for the empty U-tube one filled with fragments of pumice impregnated with perfectly anhydrous sulphate of copper. This will intercept any hydrochloric acid, and will also serve to condense any sulphuric acid vapour that may go over from the heated acid. The calculation is very simple. Every one equivalent of sodium carbonate equals one equivalent of hydro-fluosilicic acid or six of fluorine, hence the proportion—

Mol. wt. Na_2CO_3 : Mol. wt. 6F. Amt. of Na_2CO_3 : Amt. of F.
106 : 114 : : x

Below will be found some results which I have obtained from pure fluor-spar. The time of conducting the operation in every case was two hours, the amount of aspiration varied from $4\frac{1}{2}$ to $6\frac{1}{2}$ litres, and the indicator used was cochineal.

The first six were titrated with a dilute solution of ammonia, 1 c.c. of which equalled 0.0242 grm. of fluorine, the last four with a solution twice as strong and one-half of whose volume was alcohol.

Fluor-spar.	Corresponding Weight of Fluorine.	Fluorine found.	Gain or Loss.	C.c. of Air Aspirated.
0.2464	0.1200	0.1199	-0.0001	4500
0.1804	0.0879	0.0895	+0.0016	5000
0.2039	0.0993	0.0989	-0.0004	5500
0.2384	0.1161	0.1169	+0.0008	5000
0.2074	0.1010	0.1023	+0.0013	6500
0.2147	0.1046	0.1064	+0.0018	5500
0.9292	0.4325	0.4329	+0.0004	5500
0.4673	0.2277	0.2283	+0.0006	6500
0.9901	0.4824	0.4817	-0.0007	6000
1.0130	0.4935	0.4923	-0.0012	6500

—American Chemical Journal.

March 3, 1879.

SANITARY CONDITION OF AIR IN PUBLIC SCHOOLS.

By N. T. LUPTON, Vanderbilt University.

THE following series of determinations of carbon dioxide in air from the public schools of Nashville, Tennessee, was made at the request of the City Board of Health. The subject being one of general scientific interest, the results are submitted as a contribution to sanitary chemistry. (See Table on next page.)

The above observations were made each day at from 12.30 to 1.30 p.m., and from an hour and a half to two hours after recess. In a few instances, as indicated, two bottles of air were taken at the same time from one and the same room, but at different parts of the room. The results are connected by brackets. In making the determinations of carbon dioxide the well-known method of Pettenkofer was adopted, as it can be easily and rapidly performed, and gives accurate results.

Four thin white glass bottles, one of 4950, two of 3950, and one of 3925 c.c. capacity, were used. The bottles were filled with air from about the middle of the rooms while the schools were in session, by means of a small hand-bellows, and after standing a few minutes to equalise temperature and pressure, were closed with well-fitting caoutchouc stoppers. They were then taken to the open air, and exactly 50 c.c. of a solution of barium hydroxide rapidly run into each, the bottles closed, carried to the laboratory, and allowed to remain two or three hours with occasional shaking; or, rather, the solution was moved

about so as to moisten the interior surface of the and thus hasten the absorption of carbon dioxide. The milky-looking liquid was transferred to small mouthed, glass-stoppered bottles, and after the carbonate had settled, 25 c.c. of the supernatant were carefully titrated with a known solution of acid, turmeric paper being used to determine point of neutralisation. The solution of oxalic made of such strength that 1 c.c. corresponded to 10 of carbon dioxide.

For calculating the volume of carbon dioxide in volumes of air the following formula was used—

$$x = \frac{760(1 + 0.003665 \times t)p}{H(V - v)0.0019714}$$

in which H represents the height of the barometer in millimetres, V the capacity of the flask, v the volume of barium hydroxide solution used for absorption, t the temperature expressed in degrees of the Centigrade scale, p the amount in milligrams of carbon dioxide in the air experimented upon.

It will be seen from the tabular statement that the amount of carbon dioxide found in several instances as much as 25 volumes in 10,000, or one-quarter per cent. Whether this amount is sufficient to render air decidedly unwholesome may be questioned, though all will admit that an increase of six times the normal quantity found in the atmosphere shows imperfect ventilation. It was observed that in every case where the amount of carbon dioxide reached 20 in 10,000 or one-fifth of one per cent, the odour from other impurities of respiration was exceedingly offensive; and even where the amount was 10 in 10,000, offensive odour was distinctly perceptible.

While no reliable method has yet been devised for the direct determination of organic impurities resulting from respiration, recognisable by the sense of smell, and which vary with the amount of carbon dioxide in the air, and may thus be indirectly estimated.

In the construction of these school buildings, provision whatever was made for ventilation other than doors and windows, except some narrow flues in the buildings, and a ventilating-shaft at each of the corners of one other, with openings entirely too small to be of much service. The General Superintendent of schools, appreciating the importance of ventilation, established a rule which requires the teachers to leave the upper sash of each window a few inches open in the room opposite to that from which the wind may be blowing, so as to avoid draughts as much as possible during recess the windows are required to be raised, and the doors to be thrown open. It was evident from visits that these rules were faithfully carried out by the teachers, and that where defective ventilation existed, it was caused by defective construction of the room or overcrowding of pupils. It is a question of grave importance whether the return of children overheated by a cold room for the resumption of their studies is more dangerous to health than defective ventilation.

Opinions differ in regard to the quantity of air required per minute for one person for healthful respiration. Arnott says that 20 cubic feet are necessary; E. 10. Roscoe, in discussing certain experiments in school-rooms, says—"I have come to the conclusion that 10 cubic feet per minute per head is insufficient to move completely the organic putrescent matter in the air, calculating the amount of ventilation he uses the following expression:—

$$V^1 = V + Va + Va^2 + \dots + Va^n.$$

In which V^1 represents the volume of air to be supplied to produce a given amount of impurity, V the volume of air, and a the ratio between the natural impurity in the air (0.04) and that found in the room under examination. Roscoe determines only the first three terms of the

Carbonic Acid in 10,000 Volumes of Air.

Date.	Barometer.	Thermometer.		CO ₂ .	Weather	Number of Pupils in Room.	Capacity of Room in cubic feet.	Per cent of Moisture.	Cubic ft. of Air per minute for One Person.
		Dry.	Wet.						
1878.									
February 1	750	19°	16°	32.42	Cloudy	136	26,415	73	4.0
	750	18½	15½	29.29	Cloudy	137	37,377	70	4.5
6	755	19	14	15.83	Clear	136	26,415	58	9.5
	754½	20	14	15.83	Clear	137	27,377	50	13.8
8	740½	19	17	9.14	Cloudy	136	26,415	80	20.5
	740	20	17	9.71	Cloudy	137	37,377	70	23.3
11	757	15	13	8.61	Slight snow	50	12,360	80	5.0
	757	18	14½	27.22	Slight snow	50	12,360	60	6.1
12	754½	19	15	22.28	Clear	66	13,180	64	9.0
	754	19	14	16.88	Clear	50	12,360	59	5.7
13	750	18	13½	18.04	Drizzling	66	13,180	59	8.6
	749	18	15	19.31	Drizzling	50	12,360	72	10.0
	748½	18	15	14.92	Drizzling	50	12,360	72	6.0
	748	18	15	22.54	Drizzling	98	17,680	72	20.1
14	744½	19½	17	9.30	Cloudy	55	10,390	76	9.9
	744	20	18	14.84	Cloudy	55	10,390	81	7.2
	744½	19	16	19.11	Cloudy	55	10,390	73	10.0
	744	19	16	15.37	Cloudy	75	20,150	73	16.0
15	749½	19	16	11.08	Drizzling	50	10,390	73	6.6
	749	20	17	21.33	Drizzling	55	10,390	70	4.4
18	748½	19½	15	29.75	Drizzling	85	20,150	60	8.4
	757	16	11	29.50	Clear	364	61,660	54	18.0
	756½	20	16	17.50	Clear	294	61,974	64	6.0
	756	21½	17½	9.86	Clear	294	69,138	57	4.6
19	753	18	11½	28.25	Clear	364	61,660	48	12.2
	752	20½	15	26.84	Clear	294	61,974	45	5.0
	751½	20	15½	13.42	Clear	294	69,138	58	9.3
20	748	19	17	16.14	Rain	364	61,660	80	12.0
	747½	21	17	13.79	Rain	294	61,974	63	9.5
	747	21	16½	15.10	Rain	294	69,138	61	8.3
22	747	20½	16½	17.68	Cloudy	130	30,050	62	17.4
25	761	20	15½	14.18	Clear	130	30,050	58	7.8
22	748	20	16	18.58	Cloudy	125	33,850	64	9.0
25	761½	18	12	16.54	Clear	125	33,850	48	20.0
28	755	20½	14	9.14	Clear	85	15,326	49	15.6
				10.97					
				14.69					

but as it is an infinite decreasing series its exact value is—

$$V' = V \left(\frac{1}{1-a} \right).$$

To determine, then, the quantity of air in its normal condition which must be added to one of the rooms containing 136 pupils so as to give in one hour and a half the percentage of carbon dioxide (0.3242) found, we proceed as follows;—If one person exhales 0.686 cubic feet of this gas in one hour, 136 persons will exhale 139.944 cubic feet in one hour and a half. By a simple proportion we get the value of V, which, multiplied by the fraction in the equation whose terms are known, we get within reasonable limits the quantity of air with which 139.944 cubic feet of carbon dioxide must be mixed in order to make the percentage of this gas amount to 0.3242, the quantity actually found in the room. The calculation gives 49.242 cubic feet of air as necessary to reduce 139.944 cubic feet of carbon dioxide to 0.3242 of the total bulk. This allows to each person 4 cubic feet per minute, which is not sufficient for healthful respiration.

From the column under "Cubic Feet of Air per Minute for One Person," which was calculated in the above manner, it will be seen that six of the rooms visited gave only 5 cubic feet or less of pure air per minute to each

person, a quantity insufficient for healthful occupancy according to all authorities. Eleven other rooms gave only 10 cubic feet or less per minute, an amount insufficient to remove the disagreeable odours of animal effluvia. In a few instances the temperature and percentage of moisture found below the generally recognised standard, but these may have been accidental.

The general management of these schools is worthy of praise. The systematic arrangement of studies, thoroughness of instruction, and excellence of discipline, are such as to enlist the sympathy of all classes and command the patronage of those who have children to educate; but the adoption of a more efficient system of ventilation is demanded in view of the facts presented and the important interests involved.

The Discussion on Mr. Hollway's New Process in Metallurgy.—We have been requested to announce that a renewal of the discussion on Mr. Hollway's paper, "A New Application of Rapid Oxidation by which Sulphides are Utilised as Fuel," will take place at the Society of Arts on Wednesday evening next, April 30, at 8 o'clock. Dr. Roscoe, F.R.S., will preside.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, April 17, 1879.

Dr. Roscoe, Vice-President, in the Chair.

AFTER the minutes of the previous meeting had been read and confirmed, the following certificates were read for the first time:—A. E. Robinson, R. Reid, and H. Appleby. During the evening the following were declared to be duly elected Fellows of the Society:—W. E. Blythe, W. A. Bradbury, A. H. Black, T. Griffiths, W. T. Gent, C. H. Hutchinson, F. A. B. Jewson, W. Johnstone, W. T. Lawson, J. L. Macmillan, W. North, J. A. Ogilvie, F. Podmore, W. Palmer, T. Palmer, G. Rait, W. Radford, W. Stone, A. W. Stokes, W. Spottiswoode, P.R.S., C. Slater, W. B. Turner, F. L. Teed, V. H. Velez, Thorp Whitaker, T. H. Walker.

The SECRETARY then read a paper "*On Heptane from Pinus Sabiana*," by T. E. THORPE. In the *Pharmaceutical Journal*, March 30, 1872, W. Wenzell, described, under the name of Abietene, a new hydrocarbon obtained by distilling the exudation of the *Pinus sabiana*, a tree indigenous to California, known locally as the nut pine, or digger's pine. To procure the exudation, the tree during winter is notched and guttered at a convenient height from the ground. The resin on distillation yields the liquid hydrocarbon. The crude oil is met with in San Francisco as an article of commerce under the names of abietene, erasine, &c., as a substitute for benzolin, for removing grease spots, &c. It is a nearly colourless mobile liquid, of a powerful aromatic smell, resembling that of oil of oranges. Wenzell contrasts its characters with those of terebene from *P. Sylvestris*. Abietene, sp. gr. 0.694, boils at 101°, dissolves but a small quantity of hydrochloric acid gas, and is but little attacked by cold nitric acid. Terebene, sp. gr. 0.840, boils at 160°, absorbs HCl with avidity, and is violently attacked by nitric acid. From a consideration of the general properties and behaviour of this hydrocarbon the author of the present paper concluded that it was likely to be a paraffin. The occurrence of a paraffin playing the part of oil of turpentine in the vegetable kingdom was hitherto unheard of, the only natural sources of this hydrocarbon (heptane) being petroleum and fossil fish-oil. The author, therefore, obtained from Mr. Wenzell 2 gallons of the abietene, and has subjected it to a most exhaustive chemical and physical examination, the details of which are contained in the paper. The crude oil is slightly contaminated with a resinoid matter, to which its smell is due. The pure oil boils at 98.42° C. at 760 m.m. It has the composition of heptane, containing 83.85 per cent C, 16.03 per cent H (C₇H₁₆ requires C 83.97, and H 16.03). Vapour density—found, 49.94; calculated, 50.07; sp. gr. at 0° 0.70057. The rate of expansion by heat has been carefully determined: its volume at the boiling-point is 1.1411. Its specific volume 162.54; refractive index for D, 1.3879; its molecular refractive energy, 56.4. Rotates in a tube 200 m.m. +6.9°. Its viscosity and surface tension were also determined. The author has compared the heptane obtained from *P. Sabiana* with the heptane from petroleum and that obtained by heating azelaic acid with baryta. The sp. gr. of the heptane from petroleum is 0.7301; that from azelaic acid has a sp. gr. of 0.700. These heptanes are believed by Schorlemmer to be identical. The author is at present engaged in an investigation of this point.

The CHAIRMAN said the Society was much indebted to Dr. Thorpe for this elaborate investigation, and for the extremely careful and able manner in which he had worked out the subject.

Dr. ARMSTRONG said that he had detected in ordinary turpentines a paraffin-like substance in small quantities,

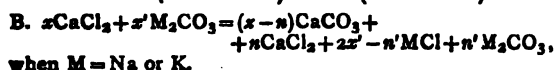
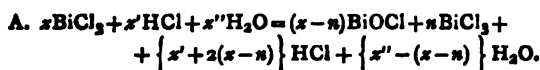
½ to 2 per cent. It was very remarkable that a paraffin should be found in such a state of purity in a plant.

Mr. ATTWOOD said that the composition of the oil would probably vary at different seasons. Sometimes the nuts of this pine tasted very strongly of turpentine, whilst at others the taste was almost imperceptible.

The next communication was "*On the Determination of Tartaric Acid in Lees and Inferior Argol, with some remarks on Filtration and Precipitation*," by B. J. GROSJEAN. "Lees" is the name given to the deposit in fermenting wine vats, "argol" being the crystalline crust which forms on the sides. The first direct method for determining the tartaric acid in "lees," &c., was suggested by Warington (*Chem. Soc. Journ.*, [2], xiii., 973); it is known as the "oxalate method." It consists in treating the finely-ground lees with a little water, and heating the mixture to 100°; an excess of neutral potassium oxalate is then added, and the whole digested for a quarter of an hour, the free acidity is nearly neutralised with caustic potash, the whole is filtered on a vacuum filter, and the residue washed; the filtrate and washings are concentrated, and an excess of citric acid added; the tartaric acid is thereby precipitated as potassium bitartrate. This salt is collected, washed, and the tartaric acid determined by titration with standard alkali. The author considers this process in detail, giving fullest details, and illustrating many obscure points with quantitative experiments:—1. *Treatment with Potassium Oxalate*.—The amount of lime present must be roughly determined (a method for this purpose is given), and 1½ grms. of the oxalate are added to ensure the decomposition of the calcium tartrate. The sample taken for analysis should contain about 2 grms. of tartaric acid. 2. *Neutralisation with Caustic Potash*.—Care must be taken to add the alkali drop by drop with constant stirring, so that the alkali is never in excess. 3. *Filtration from the Residue*. The author uses the vacuum filter suggested by Casamajor (*CHEMICAL NEWS*, vol. xxxii., p. 45). It consists of a perforated disc of platinum, lead, or pumice, placed in a funnel of peculiar shape. The disc is covered by a slightly larger circle of paper. The pressure keeps the paper so closely applied to the sides of the funnel that no precipitate passes, but at the same time a speedy and a thorough washing is ensured. The author finds that this plan has many advantages over that proposed by Bunsen, but states that an ordinary funnel can be used instead of the special funnel proposed by Casamajor. In some cases the paper disc should be covered with a layer of roughly ground pumice, freed from air, before commencing the filtration. The author recommends that the wash-water be allowed to run through until the residue presents on the surface no visible moisture, without waiting for it to crack, and then immediately to pour on a fresh portion of wash-water. The vacuum most advantageous is 250 m.m. mercury. By following this plan the potassium tartrate will be completely washed out of the sample of lees in ten washings of 2 to 3 c.c. each. This is proved by quantitative experiments. Various plans have to be adopted if the filtration refuse to proceed quickly. 4. *Addition of Potassium Chloride*.—The author recommends the addition of 5 grms. of KCl to render the precipitation of the bitartrate more complete. Thus, water at 12° dissolves 1 part of bitartrate in 262, whereas a 10 per cent solution of KCl dissolves only 1 part in 4401. 5. *Quantity of Citric Acid Required*.—An excess of 1 to 1.5 gm. of citric acid should be added. If 2 grms. of tartaric acid are present, 2 to 2.5 grms. citric acid are required to effect complete precipitation. 6. *The Mode of Precipitating the Bitartrate*.—A 50 per cent solution of citric acid is most convenient. The author prefers to precipitate the bitartrate in the state of minute granular crystals. This is effected by constantly stirring the liquid after adding the citric acid in the cold. 7. *Complete Precipitation of the Bitartrate is effected by thus stirring continuously for ten minutes*.—The author recommends that the mixture be allowed to stand for twenty minutes after the ten minutes stirring. 8. *Filtration and Washing of*

the Bitartrate.—The washing is effected by a 5 per cent solution of potassium chloride saturated with potassium bitartrate at the temperature of the air at the time of washing. 9. *The Accuracy of the Process.*—The author obtained 99.6, 99.86 instead of 100 parts of tartaric acid taken. 10. *Time Required for the Complete Estimation.*—This the author gives as four hours. In the course of his communication the author exhibited the method of filtering, the insolubility of bitartrate in a solution of potassium chloride, &c.

The next paper was entitled "*Conditions Affecting the Equilibrium of certain Chemical Systems*," by M. M. P. MUIR. The author has quantitatively studied the two reactions, which may be formulated as follows:—



In action A it is shown that the greater the value of x'' (x and n being constant) the more rapidly will equilibrium be attained; if x'' be small n will vary at different moments and will always be somewhat large. If $n=0$ the value of x varies from 450 to 460 molecules (x and x' being each equal to 1). If n is approximately equal to x , i.e., if permanent formation of BiOCl just commences, then if x' be doubled, x'' must be more than doubled, or x'' must increase in a more rapid ratio than x' . If initial equilibrium be disturbed, but without the production of a new stable state of equilibrium, the production of this disturbance is dependent not only on the relative but also on the absolute number of BiCl₃ and H₂O molecules present. The production of a new stable equilibrium is dependent only upon the relative number of molecules. In the rush of the interacting molecules the water molecules get entangled among the hydrochloric acid molecules, the greater the number of the latter (even when the proportion of HCl to H₂O is maintained constant) the greater is the entanglement. If the system has, however, time to act and react, then the result is independent of the absolute number of the molecules provided the proportion of the HCl to H₂O molecules be constant. If water be run very cautiously into a solution of BiCl₃ in HCl until a small amount of BiOCl is formed at the junction of the two liquids and the whole be then shaken up, a greater amount of BiOCl is produced in short times than if the water is added with constant stirring. During long periods of time the result is the same in both cases. In action B a condition of stable equilibrium is attained (x and x' being each equal to 1) before the whole of the CaCl₂ is decomposed; small causes tend to disturb this equilibrium. In reaction A a small amount of chemical change seems to be accompanied with but small changes in the Entropy of the system; hence the chemically reacting system soon attains a condition of equilibrium. But in reaction B a small amount of chemical change is probably accompanied with considerable changes of Entropy, so that, if time be given, the reaction will be nearly completed. In B the influence of temperature is marked when equal molecules of the reacting bodies are used; if 2 or more molecules of alkaline carbonate be used to 1 molecule of CaCl₂, the change in a given time depends but little on temperature. With equal molecules at ordinary temperature a state of permanently stable equilibrium is attained in about sixty minutes; if the temperature be raised this point of equilibrium is passed and the change is very nearly or even quite completed. Time causes an increase in the amount of chemical change, its influence being specially marked at low temperatures, i.e., when the molecular mobility of the system is small. With equal molecules of CaCl and M₂CO₃ the change formulated is complete if somewhat high temperatures and long times be employed. If the mass of one of the reacting bodies

be increased, the change is completed in a short time and at low temperatures.

Mr. M. M. P. MUIR then read a paper on "*The Action of Aqueous Hydrochloric Acid upon Bismuthous Oxide.*" When bismuthous oxide is added to aqueous hydrochloric acid the oxide is dissolved, but a point is soon reached after which the bismuth in solution is precipitated as bismuthyl chloride (BiOCl), while the oxide added is simultaneously transformed into the same salt. Finally the whole of the bismuth is present in the form of insoluble bismuthyl chloride. The action of aqueous hydrochloric acid upon bismuthous oxide, so far as the initial and final distributions of mass are concerned, may be thus formulated—



The presence of water renders possible an action between HCl and Bi₂O₃, which action would not occur in its absence.

The three following papers were, in consequence of the lateness of the hour, taken as read:—

"*On the Action of Oxides on Salts, Part II.*," by E. J. MILLS and J. W. PRATT. This paper continues the work contained in Part I., recently communicated to the Society. The authors have examined the actions of aluminic, ferric, and stannic oxides on potassic carbonate. The weight of the carbonate was the same as in Part I., and was kept constant; the temperature was about 735°, and the time three hours. The results obtained are given in a series of tables.

"*Examination of Substances by the Time Method*," by J. B. HANNAY. In a former paper the author gave an account of his method by which the thermal dissociation of a hydrated salt may easily be followed. The results of an examination of double salts, the deportment of whose components in the free state was known, are contained in the present paper. The double salts examined were the sulphate of magnesium and zinc, 14H₂O; the double sulphate of iron and magnesium; the double sulphate of copper and magnesium. Great care was taken to ensure the purity of the salts. The conclusion at which the author has arrived is as follows:—Two hydrated salts in forming a double salt containing the normal amount of water expend one half of the affinity of the anhydrous salt for its water of crystallisation in combining with each other, showing that the formation of double salts is comparable with other forms of chemical action.

"*Preliminary Note on Certain Compounds of Naphthalen and Benzene with Antimony Trichloride, &c.*," by WATSON SMITH. While distilling a mixture of antimony trichloride and naphthalen through a red-hot tube with the object of preparing dinaphthyl, this mixture was found to contain a little water accidentally admitted. On clearing out the tube long white needles were obtained; they contained antimony and carbon, but no chlorine. Heated on platinum foil the crystals melted at a red heat, burnt, and disappeared. The author believes this body to be trinaphthyl stibine or naphthyl-oxy-stibine. On melting together a mixture of naphthalen and antimony trichloride and allowing to cool, clino-rhombic crystals were obtained, crystallising unaltered from petroleum ether. The author has obtained other crystalline compounds, and thinks it probable that by the action of the antimony potassium alloy on bromo-naphthalen or bromobenzene at high temperatures trinaphthyl-stibine and triphenyl-stibine would be obtained.

The Society then adjourned to May 1, when the following papers will be read:—"On the Volumes of Liquids at their Boiling-points Obtainable from Unit Volumes of Gases," by W. Ramsay; "On a Method of Precipitating Manganese entirely as Dioxide, and its Application to the Volumetric Determination of Manganese," by J. Pattinson; "On the Determination of Nitric Acid as Nitric Oxide by Means of its Action on Mercury," by R. Warington,

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 12, March 24, 1879.

The Slow Changes Experienced by Wines during its Preservation.—M. Berthelot.—The author has made a comparative examination of two samples of port, the one made in the year 1780, and the other forty-five years old, determining the proportion of dissolved gases, of alcohol, sugars, acids, and tartar.

Remarks on Certain Points of Crystallogenesis.—Lecoq de Boisbaudran.—Since the resistance to a change of condition is not alike for the different planes of the same crystal, its solubility must vary with its outward form. Thus a supersaturated solution of basic alum being treated at a given temperature with cubes of this salt (or with portions cut according to the cubic surfaces) will not possess the same specific gravity as in cases where the desupersaturation has been effected by contact with octahedra (or portions cut according to the octahedral surfaces). The former liquid will be more concentrated than the latter, and after it has ceased giving up matter to the cubes it will still be capable of depositing upon octahedra. Even if we consider a single system of surfaces only the principle of resistance to a change of condition leads us to recognise two unequal specific gravities for a saturated liquid at a given temperature, according as we begin with a dilute or a supersaturated solution. The solubility of any substance is therefore not sufficiently defined by the quantity contained in the liquid at a given temperature in presence of an excess of the solid substance. We must further specify the species of surfaces and the direction in which the operation has been conducted. If the desupersaturation of a solution is obtained by means of crystals having several orders of surfaces there are two possible cases:—(1.) The quantity of liquid is great in proportion to the immersed masses; the crystals then assume their most stable form, and the final specific gravity is what corresponds to this system of surfaces. (2.) The quantity of liquid is very limited; the crystals cannot in these conditions assimilate matter enough to complete their form of maximum stability, and several orders of surfaces subsist indefinitely: the final specific gravity is that which corresponds to the system of surfaces destined to disappear first if the crystals could continue to grow.

Molecular Vibrations in the Magnetic Metals during the Passage of Undulatory Currents.—M. Ader.—In all the magnetic metals the passage of an undulatory current determines in their interior molecular vibrations, which, if collected, give articulate sounds. In order that the vibrations may appear with their full intensity on the exterior of the metals it is necessary to oppose to the wires or bars a mechanical action, especially the inertia of two heavy masses at their extremities.

Ytterbia, the New Earth of M. Marignac.—L. F. Nilson.—The author has succeeded in preparing perfectly pure ytterbia, presenting no trace of an absorption ray, and having a molecular weight between 131.92 and 132.17. The erbia of all preceding authors, including M. Marignac, consists chiefly of ytterbia, to which very small quantities of erbia give a rose colouration. Ytterbia is probably Yb_2O_3 .

Scandium, a New Element.—L. F. Nilson.—The author has extracted from impure erbia a substance whose spectroscopic behaviour indicates its novelty. Its atomic weight calculated for the formula of the earth ScO is below 90.

Potassium Cyanosulphite.—A. Etard.—This compound is obtained by substituting CN for OH in hydrated sulphurous acid. It dissolves in water, cold or hot, without decomposition, evolves ammonia if boiled with a caustic alkali, and reduces salts of gold and silver.

Thermo-chemical Study on the Alkalino-earthly Sulphides.—P. Sabatier.—Not susceptible of abstraction.

Certain Alcoholic Iodides and Bromides.—J. de Montgolfier and E. Girard.—An examination of the bromides of ethyl and isopropyl, and of the chloro-bromide of ethylene.

Formation of Aurin.—Ph. de Clermont and J. Frommel.—The authors contend that carbonic acid does not intervene as such in the formation of aurin, the action being due to carbon and oxygen in a nascent state.

Presence of Lithia in Rocks and in Sea-waters.—L. Dieulafoy.—Lithia is as widely distributed as soda and potassa, and accompanies these two bases in all primordial rocks. In sea-water it may be detected in the residue from the evaporation of a single c.c.

Archives Néerlandaises des Sciences.

Tome xiii., 5me Livraison.

Decomposition of Calcium Chloride by Water.—H. C. Dibbits.—Crystalline calcium chloride ($\text{CaCl}_2 + 6\text{H}_2\text{O}$) loses in dry air, even below 10° , five molecules of water, the first four of which escape readily and the last much more slowly. At 80° the salt becomes completely anhydrous in dry air. When the salt loses all its crystalline water at a temperature not exceeding 130° no appreciable loss of hydrochloric acid takes place. Between 130° and 140° the escape of acid becomes perceptible. The more the temperature rises the more hydrochloric acid escapes, still even if gently heated over a naked flame the decomposition is so slight that not more than 0.03 per cent of the acid is lost. Even at 150° the loss is so slight that it remains entirely within the ordinary limit of error.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 12, March 20, 1879.

Works of the Liebig Extract of Meat Company at Fray Bentos.—At this establishment from 10,000 to 12,000 oxen are slaughtered yearly. In 1877, 15 million kilos. of leather, tallow, and manure were exported to Europe, whilst 1½ million kilos. of *tasajo*, or dried beef, were sent to Brazil and Cuba.

Lecture on the Phylloxera.—M. Bouchardat.—The author does not think that any remedy universally applicable can be found, nor does he entertain the hope that the Phylloxera will ultimately disappear. He likens the case to that of the *Oidium Tuckeri*, a previous vine pest, and to the *Botrytis infestans*, the cause of the potato blight.

Cases of Phosphorescence Observed.—M. Nueesch.—In a certain butcher's shop all the meat became strongly phosphorescent and remained so as long as sound. If putrefaction set in and *Bacterium termo* made its appearance the luminous appearance ceased. None of the customers of the meat experienced any inconvenience from its use, and no similar phenomenon was traced in other butchers' shops in the neighbourhood.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 17, 1879.

Monohydrated Sulphate of Soda and Dihydrated Carbonate of Soda.—Julius Thomsen.—The salt deposited on heating a solution of ordinary 10-hydrated sulphate of soda, saturated at 30° was generally supposed to be anhydrous. The author finds that it retains 1 mol. of water. The salt deposited on heating 10-hydrated carbonate of soda contains not 1 mol. water but 2,

Composition of Copper Sulphide Formed in the Moist Way.—Julius Thomsen.—The precipitate formed in solutions of copper oxide by means of hydrogen or sodium sulphide is not CuS but a mixture of sulphur with a lower copper sulphide. The affinity of copper for sulphur is satisfied with the formation of Cu_2S , which is accompanied by a liberation of heat to the extent of 20,240 cal., and the reception of a further quantity of sulphur takes place without noticeable rise of temperature.

Zinc Sulph-hydrate.—Julius Thomsen.—If a zinc sulphate solution is mixed with a double equivalent of a solution of sodium sulph-hydrate no precipitate is obtained but a clear or slightly opalescent solution.

Density and Décomposition of the Vapour of Hyponitric Acid at Various Pressures below its Boiling-Point.—Alex. Naumann.—This paper consists essentially of tables of the density of hyponitric vapour at various temperatures and pressures, showing the amount of decomposition produced. It appears that at equal temperatures and with decreasing pressure the decomposition increases as also at equal pressure with increasing temperature.

Electrolytic Determination of Cadmium.—E. J. Smith.—The author has tried the aqueous solution of cadmic acetate with perfectly satisfactory results. The time required for the deposition of the metal is three to four hours. A tolerably strong current should be employed and the solution should contain 2 per cent of the metal.

Constitution of the Propyl Group in Cymol.—O. Jacobsen.

Constitution of Oxy-mesitylenic Acid.—O. Jacobsen.—These two papers are not adapted for useful abstraction.

Behaviour of Certain Nitro-Compounds with Sulphuretted Hydrogen.—F. Beilstein and A. Kurbatow.—The authors have examined chlor-dinitro-benzol, nitro-para-dichlor-benzol, chlor-ortho-dinitro-benzol, and symmetrical nitro-meta-dichlor-benzol. They state that sulphuretted hydrogen exerts a reducing action upon chlor-nitro-derivatives only when the nitro-group is not situated along with chlorine or another nitro-group.

Convenient Method for the Preparation of Mono-chlor-hydrin-sulphuric Acid.—H. Beckurts and R. Otto.

Method of Action of Mono-chlor-hydrin-sulphuric Acid.—H. Beckurts and R. Otto.

Synthesis of Aromatic Sulphones from the Chlor-anhydrides of the Sulphonic Acids and Hydro-carbons by Means of Aluminic Chloride.—These three papers do not admit of useful abstraction.

Further Contributions to the Knowledge of the Formation and Constitution of the Disulphoxides of Benzol and Toluol.—C. Pauly and R. Otto.—The authors have formerly shown that benzol-disulphoxide and benzol-zinc-mercaptid act upon each other readily and completely with the formation of benzol-disulphide and benzol-sulphinat of zinc. An analogous decomposition takes place with the corresponding toluol compounds.

Decomposition of Ethyl-disulphoxide by Potassium Hydroxide.—C. Pauly and R. Otto.—Ethyl-disulphoxide may be regarded as a thio-ether.

Action of Mono-chlor-hydrin-sulphuric Acid upon Sulpho-benzid.—R. Otto and A. Knoll.—In this preliminary communication the authors announce the formation of sulpho-benzid-sulphonic acid, with the examination of which they are engaged.

Diethyl-glyoxylic Ether and Diethyl-glyoxylic Amide.—A. Geuther.—With reference to a paper by Finnet the author points out that the two compounds above mentioned were prepared by Schreiber from dichlor-acetic acid eight years ago.

MISCELLANEOUS.

Lectures on Hygiene and Public Health.—The first lecture of Professor Corfield's course on Hygiene and Public Health will be delivered at University College on Thursday, May 1, at 4 p.m. The course will be illustrated by models and specimens from the Parkes Museum of Hygiene. Practical instruction in the methods of analysis of air, water, foods, and drugs will be given in the Hygiene Laboratory by Professor Corfield and Mr. C. E. Cassal, F.C.S., the Demonstrator.

Society of Arts.—The third course of "Cantor Lectures" for the present session of the Society of Arts is by Mr. W. H. Preece, the Electrical Engineer of the Post-Office, on "Recent Advances on Telegraphy." The course commenced on April 21. The first lecture dealt with definitions, electrical effects, and sources; economical distribution of electric currents. The second will treat principally of the transference of electricity, and will include also such subjects as wires, insulators, supports, gutta-percha, india-rubber, underground wires and cables; The third will be devoted to simple telegraphy, visual and aural signals, telephones, and telegraphic writing. The fourth will deal with duplex, quadruplex, multiplex, and harmonic telegraphy. The fifth and last is to be devoted to automatic and fast-speed telegraphy.

NOTES AND QUERIES.

Composition of Honey.—In reply to E. H. S.'s query he will find some recent investigations as to the chemical composition of honey by Dr. Campbell Brown in *The Analyst*, vol. iii., p. 267, 1878.—G. W. W.

Solvent for Gelatine.—What is the most perfect (volatile) solvent for gelatine? What work is there where the subject is fully entered into?—M. NEWMAN.

MEETINGS FOR THE WEEK.

- MONDAY, 28th.**—Medical, 8.30.
— Society of Arts, 8. "Recent Advances in Telegraphy," by W. H. Preece. (Cantor Lectures.)
— Royal Geographical, 8.30.
TUESDAY, 29th.—Civil Engineers, 8.
— Royal Institution, 3. "Mendelssohn," by Mr. Ernst Pauer.
— Anthropological, 8.
— Zoological, 1. (Anniversary).
— Society of Arts, 8. "Light Railways for Opening up a Trade with Central Africa," by John B. Fell.
— "The Advantage of Railway Communication in Africa, as Compared with any other Mode of Transport," by J. Conyers Morrell. (African Section.)
WEDNESDAY, 30th.—Society of Arts, 8. Renewed Discussion on Mr. John Hollway's Paper on "A New Process in Metallurgy."
— Geological, 6.
THURSDAY, May 1st.—Royal, 8.30.
— Royal Institution, 2. Annual Meeting.
— Chemical, 8. "On the Volumes of Liquids at their Boiling-Points Obtainable from Unit Volumes of Gases," by Dr. W. Ramsay.
— "On a Method of Precipitating Manganese Entirely as Dioxide, and its Application to the Volumetric Determination of Manganese," by J. Pattinson. "On the Determination of Nitric Acid as Nitric Oxide by Means of its Action on Mercury," by R. Warington.
— Royal Society Club, 6.30.
FRIDAY, 2nd.—Royal Institution, 9. "Action of Anæsthetics," by Prof. McKendrick.
— Geologists' Association, 8.
— Society of Arts, 8. "The Wild Silks of India, especially Tusseh," by Thomas Wardle. (Indian Section.)
SATURDAY, 3rd.—Royal Institution, 3. "Architecture," by Mr. H. H. Statham.

ERRATUM.—No. 1012, p. 170, line 12 from top, read p. 148 for p. 198.

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WILLIAM MARRIOTT

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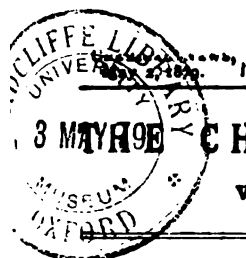
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CHEMICAL NEWS.

Vol. XXXIX. No. 1014.

RECENT CONTRIBUTIONS TO THE HISTORY OF DETONATING AGENTS.*

By Professor ABEL, C.B. F.R.S.

(Continued from p. 179.)

THE physical character of explosive substances, as also the mechanical condition of a mass of the particular explosive substance operated on, are of great influence in determining its behaviour when submitted to the action of an initiative detonation. The liquid nitro-glycerin is far more sensitive to detonation than gun-cotton; one grain of mercuric fulminate, confined in a metal case, suffices to detonate nitro-glycerin when surrounded by it; but, in order to attain this result with any degree of certainty, it is necessary so to confine the nitro-glycerin as to prevent its yielding to the blow developed by the initiative detonation, and thus to some extent escaping from the operation of the sudden concussion to which the particles contiguous to the fulminate charge are submitted.

If nitro-glycerin be mixed with solid substances in a fine state of division plastic mixtures may be obtained, and the liquid may thus be presented in something like a solid form to the detonating agent. If the particles of absorbent material be, moreover, of porous nature, as is the case with the infusorial earth called Kieselguhr, used in the production of dynamite, a solid nitro-glycerin preparation may be obtained which contains a very large proportion of the liquid (75 per cent by weight). In this condition nitro-glycerin may be detonated without any difficulty when freely exposed to air; and although it is diluted with a considerable proportion of absolutely inert material, its sensitiveness to detonation is not in the least diminished. Each particle of the diluent is enveloped in the liquid, so that no portion of the latter becomes isolated from the remainder by the admixture of inert solid matter; hence, when the initiative detonator is surrounded by such a mass it is in contact at all points with some portion of the nitro-glycerin, and the latter is in continuous connection throughout, though no longer in a mobile condition; detonation is consequently as readily established and transmitted through the mass as though it consisted entirely of nitro-glycerin. Indeed, while the liquid in its undiluted state, if freely exposed to air in a long layer, transmits detonation with difficulty and very slowly as compared with compressed gun-cotton (the observed rate of progression being, in several experiments, below 6000 feet per second), detonation is transmitted with ease and certainty through very long trains of a solid preparation of nitro-glycerin, such as dynamite, and the rate of transmission is decidedly more rapid than it is with compressed gun-cotton, a result which is in harmony with the greater sensitiveness to detonation and the greater violence of action of nitro-glycerin.

It has already been stated that gun-cotton may be detonated if a confined charge of not less than 2 grains of mercuric fulminate be detonated when closely surrounded by the substance. But in order to attain this result, the cellulose product must be presented to the detonating agent in a mechanical condition favourable to its action.

Gun-cotton in a loose flocculent condition, or even if in the more compact form of a spun yarn or thread, cannot be detonated through the agency of a large fulminate buried in the material. The light and loose gun-cotton is simply scattered with violence; portions are sometimes

inflamed by the heat developed where the fulminate is detonated, a result which is obtained with greater certainty the less violent the detonation produced by the fulminate charge. If, however, the gun-cotton be converted into a compact form, either by ramming the wool or thread very tightly into a case, or better still, by reducing the gun-cotton fibre to a very fine state of division, and compressing it, when in that condition, into compact masses, it becomes susceptible of detonation by the initiative action of mercuric fulminate, and the quantity of the latter required to bring about detonation is small (down to the limit which has been named above) in proportion as the compactness or density of the compressed material is increased.

Detonation, when established in compressed gun-cotton, is transmitted with great velocity throughout the mass, as already stated, or from one to another of contiguous masses, laid out in long rows, and even, though at a reduced rate, if small spaces exist between the individual masses. But, if a small mass of compressed gun-cotton freely exposed to air be detonated when in immediate contact with gun-cotton wool or loosely twisted yarn, the detonation will not be transmitted to these, but they will merely be scattered and perhaps inflamed.

The difference in the behaviour of nitro-glycerin and of gun-cotton when presented to the action of a so-called initiative detonation under the different conditions spoken of above, admits of ready explanation.

It was established, in the first instance, that the action of an initiative detonation is not ascribable to the heat developed within the detonating material itself in undergoing chemical metamorphosis. If it were so, the detonating mixture known as percussion-cap composition and other explosive mixtures, the detonation of which is attended by much greater development of heat than is obtained by the action of pure mercuric fulminate, should detonate gun-cotton more readily than the latter does, whereas very much larger quantities of such materials are required to attain that result. Moreover, the readiness with which gun-cotton is detonated should be solely proportionate to the amount of fulminate used, which has been shown not to be the case; and gun-cotton should be more readily detonated when in the loose and open condition than in the highly compressed or compact form, because the latter presents it in the condition least favourable, and the former in that most favourable, to ready and rapid transformation by heat. Again, the actual temperature required for the explosion of nitro-glycerin is very considerably above the exploding temperature of gun-cotton, yet a very much smaller charge is required for the detonation of nitro-glycerin than is needed for the detonation of gun-cotton. On the other hand, a quantity of confined percussion-cap composition which, if it were pure mercuric fulminate, would be altogether inadequate for the detonation of gun-cotton, suffices for the detonation of nitro-glycerin.

The action of an initiative detonation has already been compared to that of a blow from a hammer or falling weight. The readiness and certainty with which gunpowder, gun-cotton, and other explosive agents are detonated by the latter agency are regulated by several circumstances; they are in direct proportion to the weight of the falling body, to the height of its fall, and to the force with which it is impelled downwards; to the velocity of its motion; to the mass and rigidity or hardness of the support upon which the substance to be detonated rests; lastly, to the quantity and mechanical condition of the explosive agent struck, and to its sensitiveness.

Gunpowder is much more readily detonated by a sharp blow from a small hammer than by the simple fall of a heavy hammer, or by a comparatively weak blow from the latter. It is very difficult by repeated blows, applied at very brief intervals, to detonate gun-cotton if placed upon a support of wood or lead, both of which materials yield to a blow, the force applied by that blow being transferred through the explosive agent and absorbed in work done upon the material composing the support. But if the latter

* Abstract of a Paper read before the Royal Institution of Great Britain, Friday, March 21, 1879.

be of iron, which does not yield permanently to the blow of the hammer, the detonation of those substances is easily accomplished. If the quantity of the explosive agent employed be so considerable as to form a thick layer between the hammer and support, the force applied is to so great an extent expended in imparting motion to the particles of the compressible mass, that there remains little or none by which its detonation can be accomplished, and if the material be in a loose or porous condition (as in the case of a powder or a loose wool), much work has to be accomplished in moving particles of the mass through a comparatively considerable space, in the operation of compressing them, so that a second or even a third blow is required for their detonation; whereas if, by blows or pressure previously applied, the explosive material will be presented in the form of a compact mass, the particles of which have little tendency to motion when force is applied to them, detonation will be much more readily developed. It appears, therefore, that the detonation of an explosive substance by means of a blow is the result of the development of heat sufficient to bring about most energetic chemical action, or change, by expenditure of force in the compression of the material, or by establishing violent friction between its particles, consequent upon the motion momentarily imparted to them, and that it is brought about with readiness proportionate to the resistance which they oppose to their motion by the degree of their contiguity to each other.

The exceedingly violent motion of particles resulting from the sudden or extremely rapid transformation of a solid or liquid explosive body into highly heated gas or vapour (which is the effect of a detonation), must obviously exert force which operates upon a body opposed to it in a manner precisely similar to the force applied by opposing a body in the path of a solid mass which is set into very rapid motion. In other words, a detonation exerts a mechanical effect upon resisting bodies precisely similar to that of a blow from a hammer or from a projectile propelled from a gun. Just as the force of a sufficiently sudden or powerful blow from a hammer is transformed into heat by the resistance to the motion of the hammer which the particles of an opposing body offer, and by the consequent friction established between them, so the force or concussive action exerted by the matter set in motion when a solid or liquid is converted into gas or vapour, will also be transformed into heat, the development of which in an opposing body will be proportionate to the resistance to motion which its particles offer, and to the suddenness and violence of the concussion to which it is subjected. The power of accomplishing the detonation of nitro-glycerine, gun-cotton, or other highly explosive substances, freely exposed to the air, through the agency of detonation produced in their vicinity or in close contact with them, appears therefore correctly ascribable to the heat suddenly developed in some portion of the mass by the mechanical effect, or blow exerted by that detonation, and is regulated by the violence and suddenness (either singly or combined) of the detonation, by the extent to which the particles composing the mass of the explosive material are in a condition to oppose resistance to the force, and by the degree of sensitiveness of the substance to detonation, or to sudden metamorphosis, under the influence of heat thus developed.

(To be continued.)

Eupitton and Pittakall.—A. Grätzell.—The author points out a number of characteristics which distinguish eupitton from pittakall. The latter gives a blue precipitate with alkalis even in an acid solution, whilst eupitton forms a brown precipitate. Pittakall is precipitated from its acetic solution by acid acetate of alumina, whilst eupitton is not. It does not seem probable that Reichenbach was acquainted with eupitton.—*Berichte der Deutsch. Chem. Gesell.*

END-ON ILLUMINATION IN PRIVATE SPECTROSCOPY, AND ITS APPLICATIONS TO BOTH BLOWPIPE FLAMES AND ELECTRIC ILLUMINATED GAS-VACUUM TUBES.*

By PIAZZI SMYTH,
Astronomer Royal for Scotland, and Past President of R.S.S. Arts.

(Continued from p. 163).

PART III.

Electric-Spark Spectroscopy in Vacuum Tubes.

THE next step in spectroscopy, after flame lines, can hardly be anything else than the simple induction spark of electricity in rarefied gas tubes. For the most infinitely small weight of the particles to be moved or heated there, allows a very trifling quantity of electricity, and in uncondensed form, to produce abundantly visible effects. These effects, moreover, supplement in a remarkable manner our knowledge of gases as given by flame experiments. Thus on burning pure hydrogen flamewise in the open air, no spectral effects of lines or bands are seen; because, as it is said, the very act of burning being a combination of that hydrogen with the oxygen of the air, water is formed, and hydrogen ceases to exist as an independent entity. But in a close glass tube filled with hydrogen, the electric spark passing through it, heats that gas by itself alone; and then, not only do we see the hydrogen's spectral lines, but recognise them immediately as some of the most extensively distributed and most important throughout all the universe. For when the populace is in ecstasies at a total solar eclipse, on seeing the red-prominences outside the sun's circumference, and wonders what they are,—the spectroscopist pronounces them to be "flames of incandescent hydrogen of most enormous size."

Again, when the astronomer finds occasionally what is called a new star, or rather an old one which has suddenly become 500 times brighter than before, and must be a sun on fire,—a sun destroying all the life on the planets immediately around it,—it is an immense extravasation of incandescent hydrogen which has done the business. And then we are induced to speculate on whether the similar in kind, though happily smaller in quantity, extravasations, which make the red-prominence round our own sun sometimes large and sometimes small, are, or may be, regulated by cyclical law; and what length of time, if any, may probably separate us from the next of their greater effluxes?

These spectral lines of incandescent hydrogen (four in number, red, glaucous, violet, and a very faint one in the lavender) are therefore rather eerie things to look at; should never be forgotten by man; and may, in the meanwhile, be turned into, and have been already extensively used as, most useful companions at the spectroscopist, through means of the following five inventions, none of them very old.

1st. Prof. Geissler of Bonn introduced the exceeding rarefaction of gases within hermetically sealed glass tubes, armed at each end with platinum wire electrodes.

2nd. Prof. Plucker, also of Bonn, brightened up the light of those tubes by reducing the diameter of the bore of the central part until it was no more than capillary.

3rd. M. Ruhmkorff of Paris improved and vastly reinforced the induction coil.

4th. Dr. Leeson planned the bichromate galvanic battery, with its notable power, and absence of poisonous fumes; and—

5th. Some artist in arrangement produced the long-necked bottle-form of that battery, enabling its power to be called forth at intervals of either minutes, or months, or years, without any intermediate preparation or attention. For mere reference to the first three, and brightest

hydrogen lines, a one-pint bichromate bottle, with a zinc plate 3.0 by 1.5 inches, a small coil giving a $\frac{1}{4}$ -inch spark, and a transverse view are enough, and even more than enough. But for original investigations into the more untoward of the gases for lighting up, far more power and illumination are necessary. To a certain extent, any wealthy observer can obtain that higher degree of illumination most easily, by merely increasing the size of his batteries and coils with the consequent length of their sparks. But then their at-the-same-time growing heat will crack and destroy the tubes so rapidly, that even the richest man may be at last driven to inquire, whether the end-on principle just brought forward, with the claim of increasing brightness without altering the temperature at all, be not applicable?

It is a case for it undoubtedly; for the line of light in a Geissler-Plucker tube is still longer in proportion to its breadth than any blowpipe flame, and usually wastes its precious light in the proportion of 99 to 1. But as these tubes are now made they cannot be employed end-on. Their poles come in the way, two lions at once, totally obscure the view. A very considerable alteration of arrangement is therefore necessary; and not every manufacturer will take the trouble of experimenting for a single private observer. I wrote therefore to my trusty old friend M. Salleron in Paris; and he, with the enthusiasm of a scientist, joined to the practical capacities of a maker of instruments of precision, promptly realised for me one design after another of the end-on gas vacuum tubes, until we at length reached a very promising standard both of size and shape, as thus:—

The new tubes are shorter, broader, stronger than the old ones; more easily manipulated, and safely packed and carried; while, finally, in place of a uselessly long line of faint light, they give a small point of brilliant light. And they are likely to show this for a longer time; as, instead of that point of light (really the axis of the long capillary seen end-on) being looked at through the thick glass walls of the said capillary, which frequently get dimmed inside by oxidising and carbonising effect of the electric heat, they are viewed through the thin walls of the bulbs, whose very large diameter keeps down the heat on themselves and prevents those darkening effects. At the same time all the rest of the glass, except one peep-hole, being silvered externally (and then black-varnished for protection), there is some further increase of brightness by internal reflection.

I show now a box of these tubes whose light, under a standard spark of 1 inch long, may readily be pronounced by eye-view alone, extremely dull if seen *transversely*; but remarkably bright, pungent, and star-like, if looked at *end-on*. The true test is, however, the result of actual spectroscopic observation.

Viewed *transversely*, to the line of electric light, the spectrum of "air," simple atmospheric air, with a prism of 3° dispersion, begins inside the scarlet hydrogen line; and from thence to the lavender-grey, or other end of the spectrum, contains only 72 measurable features, and they of hazy, indistinct, unsatisfactory bands. But viewed *end-on*, the spectrum of the very same air tube, with the same spark and same prism, begins 18 lines or markings outside the scarlet hydrogen; and in the whole spectrum shows 221 measurable items, many of them being vividly sharp lines.

Next, on increasing the prism power to a dispersion of 22° between A and H, but everything else remaining the same, the spectral lines were found so multiplied that 400 were quickly measured in a space where our first attempt had only 38. But I do not claim that all those 400 are truly *air* spectral lines; only that they were lines distinctly seen in a well lighted tube, said to contain chemically the purest possible "air-vacuum." And in fact I stopped where I did, in the green, instead of going on to measure about 1000 more lines that were waiting in the blue and violet, because it presently appeared most indubitably, that some of the lines just entered, instead of belonging

to air in any way, were caused by carbonaceous impurities. In fact I had stumbled, and at the very right time for it on one of the grandest cases of dispute in modern spectroscopy, viz., "Is it carbon vapour, or carbon-compound vapour? carbon by itself, or carbo-hydrogen?" So I immediately increased the prism power 33° between A and H; and then, looking at the minute spectral space, which formed the crucial instance to be measured,—where, just in front of the "green giant," the best leading authority in the "Philosophical Transactions of the Royal Society," London, has only eight lines, and those dark ones,—I there saw most clearly, and measured, 31 lines, and all of them bright ones.

After such an example as that, I need not probably say anything more in furtherance of the principle of the end-on-view kind of gas vacuum tubes. But I would beg leave to call the attention of so practical a meeting as the present, to the facility of manipulating the end-on tubes by means of the peculiar holder for them when under the electric spark; also to the anti-heating character of the large, thin copper plates adopted as terminals for the induction wires, in place of the small brass wire spirals, usually employed; and to the packing-boxes lined with cork and armed with thick grooved cork; all of which have been made for me, and greatly to my satisfaction by, Mr. John Air, of the firm of Air and Candow, cabinet-makers, Leith Walk.

But the vacuum tubes themselves,* including, besides the "air-pure," examples of—

Nitrogen	Carbonic Oxide
Oxygen	Carbonic Dioxide
Ozone	Alcohol
Hydrogen	Marsh Gas
Nitrous Oxide	Olefiant Gas, and
Cyanogen	Ammonia

have all been prepared, as already intimated by M. Jules Salleron, 24 Rue Pavée au Marais, Paris; and so successfully that I do hope he may be encouraged by the patronage of many spectroscopists to apply his rare abilities more extensively to the preparation of similar gas-vacuum tubes for all known chemical volatile products. Indeed I do know that he is ready, according to the prospects which may open up, to have specially infusible glass prepared,† and to organise a system of engraving all the chief data of each tube, as it is made, on the outside of the bulbs, in a manner which must increase confidence of the scientist who uses them afterwards, exceedingly.

Coal-Tar Colours.—Mr. W. H. Perkin, F.R.S., the discoverer of mauve, the first colour produced from coal-tar, is about to read two papers before the Society of Arts on the 8th and 15th of May, in which he will give a full history of these colours, their chemistry and their technical applications.

* Experiments were made first of all as to the best area of a circular bore for the capillary; and proved that there is a *lower* limit which should not be passed if the full, as well as most intense, illumination derivable from any given electric current is to be preserved; and at the same time, there is an *upper* limit of area where the light, though abundant, becomes too faint and diffuse.

Further experiments were then made in flattening the most desirable areas of bore (as so often practised with thermometers), and adapting such flattened bore to the direction of the spectroscope's slit, in order to have a *taller* spectrum. But though the spectrum was thereby made taller, and its lines longer, the author found such bores (on the average of a dozen examples), more often clogged with opaque, adventitious particles; and after a little practice, he did not find the greater length of lines necessary to accurate micrometer observation; while in proportion to such extra length, they waste the light which might otherwise be better employed. Whence he inclines, after all, to circular bores, as more easily manufactured, and more certainly tested as to whether they correspond, or not, to any proposed general standard of size. A standard of compound *shape*, as well as size, is far more difficult to have universally kept to.

† For the ordinary tube-glass, so large an extra-dose of carbonate of soda, I am told, is usually introduced into the melting pot (to promote easy working at low heat and economise coal), that the glass becomes so hygroscopic, and depositions of both water and carbonaceous matter form in the capillaries, and can never be completely cleaned out.

EXTINGUISHING FIRES IN TAR
DISTILLERIES, &c.

By WATSON SMITH, F.C.S., F.I.C.

It may not be known to all tar distillers that in cases of fire on their premises their best remedy usually lies close at hand, and this I will shortly endeavour to demonstrate from a basis of experience and fact. In the first place, however, it may be well to state that as a rule the tar distiller cannot get any fire insurance company to insure him, and it will therefore be seen that it is a matter of no light importance to be possessed of a ready and efficacious method of subduing the dread element, when it breaks out beyond its due bounds. This remedy lies in the crude ammonia-water, otherwise known as gas-liquor, which almost every distiller works up conjointly with the tar, and if in some cases this be not so, it would be little embarrassment, and well worth while, to obtain and keep a stock of, say, 1000 gallons of this gas-liquor. The way in which I first became aware of the eminent extinguishing powers of gas-liquors (now several years ago) is as follows:—A quantity of pitch was being run from a tar-still into the pitch-house, in which it is allowed to stand for some hours, to allow most of the noisome vapours to condense. By some means a flame came in contact with the vapours, an explosion occurred, and in a moment the roof was blown off, and the whole mass of molten pitch ablaze. Water, though thrown on it in quantity, seemed to avail nothing, and at last the supply accidentally ran short. Almost in despair, the pumps for raising the gas-liquor into the ammonia-stills were adjusted so as to deliver a jet of the liquor upon the fiercely-burning pitch: the expedient acted like a charm, and the fire was quickly smothered.

I think there is considerable probability that at the high temperatures attained in some fires even the ammonia itself would be decomposed, with liberation of nitrogen and hydrogen gases, which in addition to the volumes of carbon dioxide and steam, choke out the flame by simple displacement of air. It might be said that if hydrogen and nitrogen be so liberated together, these, together with the H_2S also set free, would do much to neutralise the effect of the non-combustible gases. Not so, however, for let there be but enough incombustible gas present, as there is, to displace the air, or to choke by excessive dilution the little which may remain or diffuse, then the combustible gases present, thus for the time prevented from burning, would assist in smothering the flame (being non-supporters of combustion) with as good a will, if one may use the expression, as the incombustible ones.

I would strongly recommend every tar distiller, who also works up gas-liquor, to so arrange his pumping-gear that he may be able to throw jets of this liquor into any part of his yard or works where fire might break out and prove disastrous. If gas-liquor be not worked, then I would advise that a stock of, say, 1000 gallons be purchased. In a small works this might be advantageously stored in a tank or old boiler mounted on brickwork, at such an altitude as to give pressure enough to furnish a good jet for service below when required. Of course this reservoir should be covered to prevent evaporation.

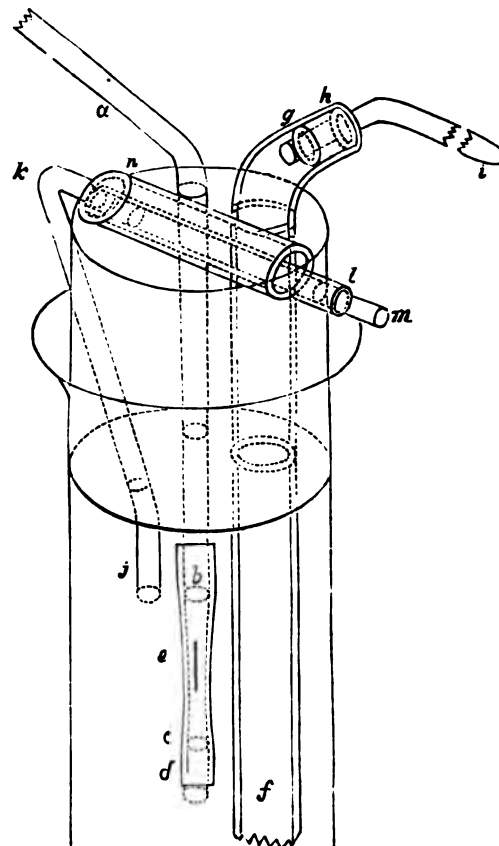
Looking at the frequent occurrence of disastrous fires in cotton-mills, especially those of Lancashire, I wrote some years ago to an eminent local paper, strongly recommending that the owners of cotton-mills should set tanks on the tops of their mills, and furnish these with supplies of ammoniacal gas-liquor, to be always ready for service. I gave, too, a short description of the best way of arranging pipes to these different rooms. The gas-liquor itself, by careful manipulation, may be pumped off very fairly clear, and free from tarry matter. Of course, besides to fires in cotton-mills and tar-distilleries, this mode of fire-extinction might be all but universally applied. It may, indeed, be readily imagined how a jet of the liquor thrown into a burning room would act, the space being so enclosed, when

the action is so powerful in a comparatively open space, and with such a refractory substance as burning pitch. It would seem, also, that the extinguishing power in the liquor would be even further called forth by a more intensely hot fire, for here, in all probability, the ammonia would be decomposed, and the generated nitrogen and hydrogen gases would help to swell the volume of non-supportive gas, displacing the air, and so choking out the combustion.

A NEW FORM OF WASH-BOTTLE.

By F. R. MALLET,
Geological Survey of India.

In the CHEMICAL NEWS, vol. xxxix., p. 19, Mr. M. H. Foye gives a description of an ingenious form of wash-bottle, the special feature of which is the introduction of a short piece of india-rubber tube in the course of the mouth-tube. By compressing this between the cork and the finger communication is cut off between the mouth and the bottle. The advantages of this are (1) that after blowing into the bottle and then compressing the rubber tube, the pressure of air is sufficient to maintain the jet of liquid for some time; this jet can be instantly stopped



by releasing the rubber tube. (2.) That steam or unpleasant gas is prevented from passing from the bottle into the mouth; and (3) that when pouring liquid out of the mouth-piece the flow can be accurately regulated by pressure on the rubber tube.

On proceeding to make a wash-bottle according to the

instructions given by Mr. Foye, it struck me that, for use with hot or unpleasant liquids, some improvement would be effected by combining Mr. Foye's plan with that of another form of bottle, which is described in Prof. Thorpe's "Quantitative Analysis" (p. 55). The special feature of this latter form consists in the attachment to the bottom of the mouth-tube (*a b*) of a valve, constructed of a piece of rubber tube (*b d*), closed at the lower end by a bit of glass rod (*c d*), and provided with a slit (*e*), which remains closed except when the mouth-piece is blown into.

The advantage of this form of bottle is that unpleasant gases are completely cut off from the mouth-piece (*a*). But there are the defects (1) that it cannot be used for hot water, because the only exit from the bottle being by the jet-piece (*f h i*), as soon as the water approaches the boiling-point the pressure of steam causes it to dribble or squirt out of the jet; and (2) that in the case of cold liquids, after the bottle has been blown into, the jet of liquid continues until the elastic force of the air inside has been expended; hence the jet is not fully under control.

In Mr. Foye's bottle the jet is completely under control. In the case of hot water, however, the mouth-piece (*a*) is the exit for escaping steam, and hence, although this steam can be cut off before applying the lips, the mouth-piece has already been rendered more or less hot. With unpleasant gas, after the rubber tube is closed, the mouth-piece above the rubber tube still contains some of this gas, which is free to enter the mouth. Forgetfulness to remove the lips before releasing the rubber tube will allow gas or steam from the bottle itself also to enter the mouth.

The modified form of bottle that I have adopted is represented in the accompanying sketch. The mouth-piece has the valve (*b d*) attached to the lower end. *f g i* is a jointed jet-piece of the ordinary form, the small glass tube (*i h*) being inserted into the large one (*f h*), with a bit of rubber tube (*g h*) between. *j k* is a third tube, penetrating the cork obliquely, and bent at *k* at an acute angle. A piece of rubber tube (*l*) is, at one end, slipped on to *k* and terminated at the other by a short bit of glass tube (*m*). Outside the rubber tube (*l*) is one of considerably larger diameter (*n*). If such a bottle be used for hot water, the steam escapes from *m*, and hence the mouth-piece (*a*) remains perfectly cool. On compressing the tube *n* with the forefinger and blowing into *a*, a jet issues from *i*, which continues after the removal of the lips until the elastic force of the air in the bottle is expended, but which can be instantly stopped by releasing *n*; the use of the outer tube (*n*) is as a non-conductor. The tube *l* becomes hot from the issue of steam, but *n*, being separated from it by an air space, remains sufficiently cool for the finger to be placed on it without discomfort. A bottle of this sort must, of course, have the neck wound round with twine or other material to keep the hand cool. If the bottle be used for cold liquid only *n* may be dispensed with.

By shifting the valve (*b d*) from the mouth-tube to the tube *j*, the bottle may, if desired, be instantly converted into one on the ordinary plan, or if this be considered unnecessary, the bend of the tube at *j* (which must be made after the tube has been inserted into the cork) may be dispensed with.

For cold water Mr. Foye's bottle is preferable, as liquid can be poured out by the mouth-piece and the flow accurately regulated, which cannot be done with that I have described.

Calcutta, March 27, 1879.

The Government Patent Bill.—A paper on the "Government Patent Bill" will be read before the Society of Arts on Wednesday evening next, the 7th inst., by W. Lloyd Wise, A.I.C.E. F. J. Bramwell, F.R.S., in the Chair.—*Ber. der Deutsch. Chem. Gesell.*

EXPLOSIONS IN FLOUR MILLS.

By H. W. LANGBECK.

IN the *Annales de Chimie et de Physique* there appeared, last year, a communication from L. Smith to Dumas concerning the explosive mixtures of finely-divided organic substances and air, especially of flour and air in some mills at Minnesota. Mr. Smith supposes that meal-dust diffused in the open air was inflamed through the heat produced by the friction of the millstones turning with great rapidity. I think I am justified in opposing that explanation and substituting another one.

Years ago I visited a friend, proprietor of a steam-mill, and there I noticed that the grains crushed between the millstones produced a smell like that of impure hydrogen. I called my friend's attention to the danger that might arise from want of sufficient ventilation, but laughingly he replied that there was nothing to fear. I forgot the matter until the above-mentioned explosion reminded me of it. In order to convince myself of the correctness of Mr. Smith's supposition, I put some meal-dust, previously heated, in a dry bladder, blew air into it, and having diffused the dust by shaking, I directed the point of a gas-flame by means of a blowpipe into the opening of the bladder (an electric spark would have been preferable, but was not at hand). No explosion took place. I allowed, then, coal-gas to enter the bladder, and repeating the experiment, a violent detonation was the expected consequence. Meal-dust diffused in a balloon, and blown through a gas-flame, was ignited; but this was, of course, not the condition on which the explosion at Minnesota took place. But what may be the source of the hydrogen, if indeed present? The amount of fat in wheat flour varies between 1 to 1.9 per cent, of bran 4 to 4.7 per cent, of maize, perfectly dried, 8 to 9 per cent. Is water predisposed to oxidise the fat through heat produced in grinding, thereby setting free hydrogen? Has the change of a small amount of starch into dextrin and glucose during the grinding process something to do with it?

CONVERSAZIONE AT THE ROYAL SOCIETY.

ON Wednesday evening last the President of the Royal Society, Mr. W. Spottiswoode, gave a Soirée at Burlington House, which was largely attended. The novelties exhibited in the various rooms attracted great interest. In the first room were exhibited Mr. Crookes's Exhausted Tubes and Apparatus illustrating various phenomena connected with Molecular Physics in High Vacua. The following is a description of the experiments which were shown:—

1. *Dark Space round the Negative Pole.*—When the spark from an induction coil is passed through an ordinary vacuum tube, a dark space is seen round the negative pole. The shape and size of this dark space do not vary with the distance separating the poles; nor, only very slightly, with alteration of battery power, or with intensity of spark. This well-known dark space appears to be a layer of molecular disturbance identical with the invisible layer of molecular pressure or stress, the investigation of which has occupied the exhibitor some years.

2. *The Electrical Radiometer.*—An ordinary radiometer is furnished with aluminium cups for vanes. The fly is supported by a hard steel cup, and the needle point on which it works is connected with a platinum terminal sealed into the glass. At the top of the radiometer bulb a second terminal is sealed in; the radiometer can therefore be connected with an induction coil, the movable fly being made the negative pole. At low exhaustions a velvety violet halo forms over each side of the cup. On increasing the exhaustion the dark space widens out, retaining almost exactly the shape of the cup; the bright

margin of the dark space becomes concentrated at the concave side of the cup to a luminous focus, and widens out at the convex side. On further exhaustion, the dark space on the convex side touches the glass, when positive rotation takes place.

3. *Green Phosphorescent Light of Molecular Impact.*—At very high exhaustions the dark space becomes so large that it fills the tube, and when German glass is used the sides are beautifully illuminated with a greenish yellow phosphorescent light.

4. *Projection of Molecular Shadows.*—The rays exciting this green phosphorescence will not turn a corner in the slightest degree, but radiate from the negative pole in straight lines, casting strong and sharply-defined shadows from objects which happen to be in their path. The best and sharpest shadows are cast by flat disks, and not by narrow-pointed poles; no green light is seen in the shadow itself, no matter how thin, or whatever may be the substance from which it is thrown.

5. *Magnetic Deflection of the Trajectory of Molecules.*—The stream of molecules, whose impact on the glass is accompanied by evolution of light, is very sensitive to magnetic influence, and the shadow can be deflected by bringing a small permanent magnet near, the amount of deflection of the stream of molecules being in proportion to the magnetic power employed. The trajectory of the molecules forming the shadow is curved when under magnetic influence.

6. *Focus of Heat of Molecular Impact.*—Great heat is evolved when the concentrated focus of molecular rays from a nearly hemispherical aluminium cup is allowed to fall on a strip of platinum-foil, the heat sometimes exceeding the melting-point of platinum.

7. *Mechanical Action of Projected Molecules.*—An actual material blow is given by the impinging molecules. A small vaned wheel being used as an indicator, by appropriate means the molecular shadow of an aluminium plate is projected on the vanes. When entirely in the shadow the indicator does not move, but when the molecular stream is deflected so that one-half of the wheel is exposed to molecular impact it rotates with extreme velocity.

8. *Phosphorogenic Properties of the Molecular Stream.*—Substances known to be phosphorescent under ordinary circumstances shine with great splendour when subjected to the negative discharge in a high vacuum.

(a.) *Becquerel's Luminous Sulphide of Calcium* shines with a bright blue-violet light, and when on a surface of several square inches is sufficient to faintly light a room.

(b.) *The Diamond* is very phosphorescent. Most diamonds from South Africa phosphoresce with a blue light. Diamonds from other localities shine with different colours, such as bright blue, apricot, pale blue, red, yellowish green, orange, and pale green. One large fluorescent diamond gives almost as much light as a candle when phosphorescing in a good vacuum.

(c.) *The Ruby* glows with a rich full red, and it is of little consequence what degree of colour the stone possesses naturally, the colour of the phosphorescence is nearly the same in all cases.

In another room Prof. Guthrie, F.R.S., exhibited Broken Glass in Frames, illustrating the Fracture of Colloids.

In Room V. great interest was taken in the working of the Writing Telegraph, exhibited by Mr. E. A. Cowper. The object attained by this instrument is that it enables the operator to write at a distant station many miles away, just as though he were present there himself, without requiring the use of any special signals, codes, or signs (to spell each letter as is now the practice), and without the assistance of any person to translate the signals as received. The instrument acts upon the simple principle of communicating at all times, to the writing pen at the receiving end of the line, the exact position of the pencil of the operator at the sending station through two line wires, or, so to speak, giving the latitude and longitude of the pencil continually, the position of the pencil vertically being communicated by one wire, and the

position horizontally being communicated by the other wire. The pencil of the operator has two light "contact rods" jointed to it, and one of these slides over the edges of a series of "contact plates," having various resistances interposed between them and the line wire, while the other rod slides over a second set of such plates connected to the other line wire; at the receiving end of the line each of these wires actuates its own needle. The two needles (which are placed at right angles to each other, and are provided with light springs) actuate one writing pen, so that the pen moves up or down, and backwards or forwards, in exact obedience to the motions of the pencil in the hand of the operator at the distant station. Both the paper written upon in pencil by the operator at the sending station and that written upon in ink by him at the receiving station move along as the writing proceeds, and the messages have only to be cut off from time to time, wound round a piece of card and sent out to their destination, or put into an envelope and dispatched.

In the Meeting-Room Edison's Loud-speaking Telephone was exhibited by Mr. Arnold White and Mr. C. P. Edison.

Messrs. Preece and Stroh exhibited a synthetic curve machine, and frame of curves produced thereby. Automatic phonograph. Electro-magnetic vowel-sounder. Stereoscopic curves. Synthetic sounder and syren. Phonautograph.

Mr. J. Browning exhibited a diffraction spectroscope. Automatic spectroscope. Mayall's automatic spectroscope with very dense glass prisms. Automatic sunlight recorder. New bisulphide of carbon prisms. New automatic electric lamp; burns equally well horizontally or vertically.

Mr. A. Hilger exhibited a quartz spectroscope for the ultra-violet rays: constructed for the Scientific Society at Stettin under direction of Dr. Schönn. Universal Christie half-prism spectroscope. New spectroscope, devised by Professors Living and Dewar. Thollon high-power dispersion bottle prism. Hilger's universal variable power prism, from 2° to 10°; collection of different power prisms and pocket spectroscopes; (A to H) dispersion.

Messrs. Tisley and Co. exhibited Tisley's dynamo-magneto machine, with single wire on armature: the alternate currents are used to augment the magnetism and for external work; the armature being hollow is kept cool by a stream of water flowing through. Donkin's harmonograph with parallel and rectangular motions.

PROCEEDINGS OF SOCIETIES.

PHYSICAL SOCIETY.

Ordinary Meeting, April 26, 1879.

Prof. W. G. ADAMS, President, in the Chair.

Mr. C. V. BOYS gave an account of some experiments made by Dr. Guthrie and himself on the subject of Arago's rotation. The experiments were begun with a view to determine if the drag on a copper disk when a magnet is made to revolve beneath it, or on the magnet if the disk is made to revolve above it, could be made use of for determining the velocity of running machinery. They made the magnet revolve and obtained the angle of deflection of a disk suspended by a torsion thread (the hair spring of a watch.) They found, as Snow Harris and others found before, that other things being equal, the drag is directly proportional to the speed, so that if the torsion of the thread could be relied on, and the strength of the magnet did not change, a perfect velocimeter could be constructed. They consider that this method is better than observing the deflection of a magnet over a revolving disk, as in this case they are limited to less than a right angle, and changes in the absolute magnetism of

the earth would affect the results. They also determined the effect of change of distance, thickness, diameter, and nature of the disk, &c., their results agreeing with those of former experimenters. They observed that the effect of concentric circular cuts was far greater than that of even many radial cuts; and that when radial sectors were entirely separated from each other the effect was much less than when these were united at the centre. They then experimented on liquids by suspending a sphere or cylinder of the liquid between the poles of a revolving electro-magnet, and succeeded in getting a decided and measurable effect. The importance of this is very great, for they have thus a means of determining the conductivity of liquid electrolytes by currents induced in the liquid without the use of electrodes and without polarisation.

Dr. GUTHRIE stated that as the push on the liquid is directly proportional to the current quantity they hope to measure the conductivities of liquids, and connect these to the conductivity of solids through the intervention of mercury.

In reply to Prof. Adams,

Mr. Boys said that the angle of deflection of the conductor had proved to be proportional to its conductivity.

Dr. O. J. LODGE suggested that the conductivity of the disks used in these experiments should be determined by plotting out the equipotential surfaces.

Dr. SYLVANUS THOMPSON recommended trying conducting jellies in these experiments, and

Dr. GUTHRIE replied that such were being prepared for trial, including the permanent jelly made by dissolving gelatine in anhydrous glycerine at 100°.

Prof. SYLVANUS THOMPSON then communicated five laboratory notes from University College, Bristol. The first related to the source of sound in the Bell telephone receiver. Two theories are now being discussed as to this effect, the molar theory regards the motion of the diaphragm in mass as the source of sound; the molecular theory finds it in the molecular motions of the magnetic core of the instrument. Prof. Thompson applied his method of getting magnetic curves with iron filings dusted on gummed glass to this problem. He found that when no currents passed in the telephone the magnetic lines springing from the pole of the magnet are gathered together on the diaphragm opposite, over a central region, which is magnetised lamellarly, or like a magnetic shell. The rim of the plate beyond this region is, however, magnetised radially, and between these two zones there is a neutral circle. It was remarkable, too, that the lines of force touching the plate were bent back around the circle, forming a kind of valley. When the current passed in the coil in a direction so as to reinforce the magnetism, the lines are gathered more closely on the central region of the plate. If the current diminishes the magnetism the lines are, on the other hand, repelled from the plate. The neutral line is also altered. In the first case it shrinks in size, in the second it expands. A small thick disk is wholly magnetised lamellarly; a disk entirely magnetised radially becomes slightly conical in shape. In the actual telephone the disk is flat at the middle and conical at the edges. As the neutral ring shifts the diaphragm will assume new nodal lines. Dr. Thompson concludes that the molecular theory is not, therefore, necessary to account for the speech of the telephone, although it may assist. As confirming this view he found that with iron rings round a cardboard diaphragm and an iron centre piece the enunciation was good, though the *timbre* was altered; whereas with radial pieces of iron on the cardboard the *timbre* was good, but the enunciation bad. In reply to Prof. Adams, Dr. Thompson said that the stronger the magnet the smaller the lamellarly magnetised space became, and that with a thicker disk the neutral ring was not so well marked.

Dr. LODGE suggested that the best place for the coil would be in the valley over the neutral ring, which was in an unstable condition.

Dr. THOMPSON next wrote on a saw blade with a magnet, and dusted iron filings on it, which arranged themselves so as to trace the writing. This is usually shown on a steel plate, but a saw retains the virtue for six or eight months. A modification of this experiment due to himself consisted in writing on the blade with one pole of a powerful battery, the other pole being connected to the end of the blade.

The third "note" recommended the use of fine steel fibres, got by breaking iron gauze of 32 meshes to the inch, instead of iron filings for exhibiting magnetic lines. The fourth note showed that the lines of force got by filings fixed on cards are magnetic, that of a magnet acting as a magnet. The fifth note explained that solid magnetic "figures" could be got by coating iron filings in shellac to make them light, and floating them in water, or by mixing filings in a soft paste of plaster-of-paris, which could be cut into sections on hardening.

CORRESPONDENCE.

NEW FORMS OF APPARATUS.

To the Editor of the Chemical News.

SIR,—Will you allow me, as a reader of your esteemed journal, the CHEMICAL NEWS, to address a few lines to you concerning an article contained in vol. xxxix., p. 160.

The article in question which attracted my attention is that of Mr. M. Benjamin, "On some New Forms of Apparatus," a water-bottle and a burette clamp. I am astonished to see these apparatus described with the predicate of new ones; as when I began my studies as a chemist in the year 1871 they were quite usual things in the eyes of my fellow students. In a great many laboratories this water-bottle is adopted; I only mention the laboratories of the Universities of Leipzig, Bonn, and that of the new Polytechnische Schule of Dresden. But this the distant describer may possibly not know; I therefore draw his attention to the *Zeitschrift für Analytische Chemie*, by Fresenius, where he will find, Vol. II., page 190, from the year 1872, this recently devised apparatus described by H. Fresenius. This description is illustrated by a woodcut, which differs from the one in the CHEMICAL NEWS by only showing the chief parts, the tube through which the water flows and fills the bath to a height, which is determined by a vertical tube through which an excess of water is allowed to flow out. The bath itself is left out in the woodcut, but this may, with very little imagination, easily be added. As to the other new apparatus spoken of in Mr. Benjamin's essay, I will only mention, in a few words, that this is just as old as the first, and even older. It is, and was, to be found in the catalogues of every business of chemical apparatus of some significance, such as Lezbold, Köln, Gehrhardt, Bonn, Noellner, Darmstadt, and I do not doubt that it also is to be found in English and American businesses.—I am, &c.,

DR. FERDINAND KORN.

Renteria, April 16, 1879.
Guizuzcoa Espana.

LOSS OF NITRE IN VITRIOL MANUFACTURE.

To the Editor of the Chemical News.

SIR,—I should have thought, from Dr. Hurter's last reply to me in *Dingler's Journal* and for other reasons, that he would not renew the discussion upon the loss of nitre, at least upon precisely the same lines as those previously taken. But as he thinks fit so to do, and this time before the English public, I am simply compelled to reply to him once more. Those who are further interested in this matter I refer to my explicit refutations of Dr. Hurter's views and

statements in *Dingler's Journal*, vol. cccxviii., pp. 70, 152, and 548.

The principal reason why I have always contended that in the Glover tower no appreciable loss of nitre takes place is this:—That notoriously since the introduction of that apparatus the consumption of nitre in vitriol works, everything else being equal, has not merely been found to be no larger than before—it has actually decreased, and, in some cases, considerably. For instance, it fell from 2.0 to 1.3 parts of nitre to 100 parts of pyrites, according to the figures furnished to me by the manager of one of the best French works, and the fact of that decrease having taken place is not disputed by Dr. Hurter himself in his German papers. But "all fear of appreciable loss of nitre in the Glover tower has vanished now," at any rate, among the general body of vitriol makers, although Dr. Hurter, and possibly a few others, take exception to that view. This I can positively affirm from numerous verbal and written communications of vitriol makers in all principal countries (whom, if need be, I could quote by name); and the fact adduced in my last letter, viz., that some of the most careful manufacturers (e.g., Schaffner of Aussig) introduce all their nitric acid through the Glover tower is a striking proof of the above. Indeed, the all but universal use of the Glover tower is the best practical refutation of the reproach cast upon it by Dr. Hurter, and before him by M. Verster. It therefore requires very weighty and accurately defined arguments to substantiate the statement that an appreciable, let alone a large, loss of nitre takes place in the Glover tower. Has Dr. Hurter produced any arguments of that kind? He estimates the loss of nitre in the manufacture of vitriol from various sources, and lays the unaccounted-for balance to the charge of the Glover tower. It is unnecessary to say how little conclusive such a reasoning process is, for the result is only a margin, arrived at indirectly, and is vitiated by all the errors in estimating the remaining possible sources of loss. It could only be admitted if the latter estimation was in all cases based upon thoroughly reliable data.

For argument's sake, and as I am extremely unwilling to open out any side issue, I will grant Dr. Hurter's statement of the "mechanical losses," amounting to 25 per cent of the nitre, or 1 part of nitre to 100 sulphur.

Coming to the "chemical losses," Dr. Hurter declines to admit any loss of nitre by the oxidation of arsenious acid in the Gay-Lussac tower. Here he is at variance, not merely with the statements of Davis, but also with the very careful investigation of H. Hjelt (*Dingler's Journal*, vol. cccxvi., p. 174). Nor can I agree with him that nitric oxide cannot escape as such, on the ground that it would be re-converted into higher oxides by the excess of oxygen present. That nitric oxide sometimes does occur in the exit-gas, even when working under normal conditions, i.e., with an excess of oxygen, is proved by the ruddy vapours which are formed at the mouth of the chimney, although the "sight" in the exit-pipe is altogether colourless. This is easily explained by the fact that there is not time for the gases to be thoroughly mixed during their rapid transit through the tower, so that in some places free oxygen, and in others NO, may be present.

But now I come to Dr. Hurter's principal fallacy. This is the assumption that the "chemical loss" in the chambers themselves does not exceed 20 per cent of the nitre introduced into the system. But this is the very point to be proved, and such a proof is not even attempted by Dr. Hurter, otherwise than by quoting from Mr. Davis, to whose paper in every other respect he denies any demonstrative power. But here he should most assuredly have maintained his denial with all jealousy; for from Mr. Davis's position it might be inferred that in factories making their acid from brimstone, and working with a nitre-recovery apparatus, next to no loss of nitre should take place, as they have not to contend with arsenic! But here I am able, as I was in my German papers, to turn Dr. Hurter's own arguments against himself. He states the "mechanical losses," and that, as he thinks, with great

exaggeration, at 25 per cent of the nitre; and he declines to admit any "chemical loss" other than that taking place in the Glover-tower or in the chambers. Consequently, in factories working without a Glover-tower, the total loss of nitre could only be 25 per cent of that assumed as the basis of his calculations, viz., 4 parts per 100 of sulphur, that is, 1 part per 100 of sulphur. To this could only be added the amount of nitre lost in the chamber-acid over and above the 10 per cent allowed for already in the above 25 per cent. Mr. Davis found 8.7 and 3.6 per cent of the nitre in undenitrated chamber-acid (leaving out the "only occasionally watched" cases). Dr. Hurter's own statements in *Dingler's Journal* are no use to us here; for they refer to works where, in the first instance, altogether too much nitre was lost (5 per cent of the sulphur), and where, secondly, the very existence of a Glover-tower and the possibility of denitrating the chamber-acid, would, quite properly, cause this to be kept rather more nitrous than where the Guy-Lussac acid is denitrated by steam, and all the nitre contained in the chamber-acid is lost. The only statements referring to the latter case upon which I can lay my hands are the following:—Kolb (*Bull. Soc. Chim.*, 1872, p. 309) states the percentage of N_2O_5 in the "large chamber" (that from which the vitriol is withdrawn for use) = 0.010 per cent, the acid containing 58.6 per cent SO_4H_2 . This is equal to 0.046 N_2O_5 , or 0.100 $NaNO_3$ upon 100 sulphur. He does not state the amount of nitre used, but we may safely assume it at least = 4 per cent of the sulphur, and thus we arrive at only 2.4 per cent of the total nitre as lost in the chamber-acid.

Hasenbach (*Ber. der Deutsch. Chem. Ges.*, vii., p. 681) arrives at a loss of 31 cwts. of nitrate for 60,000 cwts. of vitriol of 66° Baumé, which he states to be about 6 per cent of the total nitre used (it is about 0.19 nitre to 100 sulphur).

Scheurer-Kestner (*Comptes Rendus*, November, 1875), in order to save his platinum stills, keeps his chamber-acid rather sulphurous, and thus practically quite free from nitre; but he has himself stated that his consumption of nitric acid (at 36° Baumé) is 3 per cent of the pyrites, equal to about 4.7 nitre to 100 sulphur. Consequently, if we allow 10 per cent of the nitre as lost in the chamber-acid, also for works without a Glover-tower, we are very much on the right side, according to all authorities. Now this brings our total possible loss, apart from that in the chambers themselves, for works denitrating by steam, to 25 per cent, leaving 75 per cent of the loss to take place in the chambers themselves (whether by Weber's reaction or in any other way), not 20 per cent, as allowed by Dr. Hurter. Thus the big margin claimed by him as chemical loss in the Glover-tower suddenly vanishes into thin air.

I should be the last person in the world wishful to "put a manufacturer off his guard" and "prevent further search" by my statements. But I cannot certainly, on the other hand, run to the opposite extreme, and decline to accept the fact that hitherto no satisfactory proof has been given for any loss of nitre occurring in the Glover-tower, and that all unbiassed evidence goes the other way.—I am, &c.,

GEORGE LUNGE.

Technical Laboratory of the
Federal Polytechnic School, Zürich.

The Wax of *Ficus Gummiiflua*.—F. Kessel.—This wax is a chocolate-coloured mass, which softens when heated and melts between 60° and 70°. Boiling water extracts a considerable quantity of a brown colouring matter, leaving the wax nearly white. Boiling alcohol dissolves considerable quantities of the wax, which on cooling are re-deposited in a white mass resembling cauliflower. By treatment with ether the wax may be separated into two bodies more or less soluble, the former having the composition $C_{15}H_{30}O$ and the latter $C_{27}H_{54}O$. —*Ber. der Deutsch. Chem. Gesell.*

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 13, March 31, 1879.

Conformity of the Systems of Fractures Obtained Experimentally with the Systems of Joints Intersecting the Cliffs of Normandy.—M. Daubrée.—In general these joints form two systems, one of which corresponds to the direction, and the other to the line of the greatest slope.

Convenience of Special Denominations for Different Orders of Fractures of the Crust of the Earth.—M. Daubrée.—The joints, or as he suggests, disjoints, due to rupture, the author would call diaclasses. When the fracture is accompanied by displacement he terms it paraclasses, whilst as a common name for both he suggests litho-clase.

Chemical Researches on a Filamentous Matter found in the Excavations at Pompeii.—S. de Luca.—The filaments in question seem to be carbonised fibres of flax or hemp, and had probably been collected for lint.

Thermic and Galvanometric Laws of the Electric Spark Produced in Gases.—E. Villari.—The heat developed in gases by an electric spark is proportional to the quantity of electricity which produces it. The galvanometric deviations produced by the discharge of Leyden jars are proportional to the quantities of electricity condensed. The galvanometric deviations produced by one and the same charge of the condensers are constant and independent of the length of the spark produced at any given point of the circuit. If one and the same quantity of electricity amassed in any condenser is discharged through a metallic circuit, interrupted so as to give rise to a spark, the quantity of electricity brought into play in the discharge is constant and independent of the length of the spark. The quantity of heat developed in any gas by the electric spark increases in proportion to its length, whence the temperature of the spark at its different points is independent of its length, and the electric resistance of the gases is proportional to the thickness of the gaseous layer traversed by the discharge. When the charge producing the spark remains the same the quantity of heat developed by this spark is independent of the surface of the condenser. In fine, the thermic and galvanometric deviations produced, the former by the spark and the latter by the discharge of a condenser, are proportional to the quantity of electricity which produces them, and at the same time to the length of their active circuits (this name being applied either to the length of the spark or to the length of the galvanometric wire).

Magnetic Rotatory Power of Gases at Ordinary Pressure and Temperature.—H. Becquerel.—By means of a special arrangement which the author describes he has not merely succeeded in showing the magnetic rotation of the plane of polarisation of light at the ordinary temperature and pressure, but has been able to measure it with precision.

Rotatory Magnetic Power of Vapours.—E. Bichat.—On causing the current of eighty large Bunsen elements to act upon a ray of polarised light traversing vapours of carbon sulphide the author has obtained an evident but slight deviation of the plane of polarisation.

Pressure Exerted by Galvanic Deposits.—M. Bouty.—Not suitable for abstraction.

Alkaloids of the Pomegranate.—C. Tanret.—The author finds that pelletierin is accompanied by three other volatile bases, the extraction of which he describes.

Formation of Carbonic Acid, Alcohol, and Acetic Acid in the Presence and Absence of Alcohol.—A. Béchamp.—The same products are generated in either case.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 17, 1879.

Reductive Action of Milk-sugar upon Alkaline Solution of Copper.—H. Rodewald and B. Tollens.—The authors point out that whilst chemists are agreed on the reductive power of dextrose, there is great discrepancy concerning lactose, 1 mol. of which is considered to represent various quantities of copper oxide, ranging from 64 to 8 atoms. They find that the exact quantity required is 7.47 atoms of copper to 1 mol. of milk-sugar, = 6700 mg. milk-sugar to 1 c.c. of Fehling's liquor. They recommend that this reagent should not be prepared in quantities beforehand; 60 grms. of the best caustic soda and 173 grms. of re-crystallised tartrate of potash and soda are dissolved in $\frac{1}{2}$ litre of water, whilst 34.630 grms. pure copper sulphate are dissolved separately in another half litre. Equal volumes of these two liquids are mixed when wanted.

Oxidation of Xylol-sulphamids.—Ira Remsen.—A reply to a communication by O. Jacobsen, which the author characterises as "almost purely personal."

Formation of Ethylamin.—H. Köhler.—If the black powder produced by agitating calomel with ammonia is heated in a glass tube, through which a current of dry ethyl-chloride is passed, a small quantity of ethylamin is produced.

Certain Colouring Matters of the Rosanilin Group.—E. Fischer and O. Fischer.—If a solution of dimethylanilin in dilute sulphuric acid is mixed with manganese the formation of violet derivatives begins at 30° to 40°.

Meta-nitro-phenol and its Derivatives.—A. Bautlin.—An interesting paper, but not susceptible of useful abstraction.

Certain Greek Tanning Materials.—Hans Jahn.—The author states the average proportion of tannin in the cups of valonias, freed from scales at 22.615 per cent, in the scales at 36.60, and in the entire ware at 33 per cent. Peloponnesian gall-nuts contain on an average 47.6 per cent of tannin; pine-bark from Crete 9.818 per cent, and from Asia Minor 17.285 per cent.

Behaviour of Mono- and Dibrom-acetic Ethyl ethers with Aqueous Ammonia.—F. Kessel.—The author sought to use the behaviour of these bodies with aqueous ammonia as a means for their separation, but succeeded only when the dibrom-acetic ether formed the greater part of the mixture.

Etherification of Secondary Alcohols.—N. Menschutkin.—Not suitable for abstraction.

Cantharic Acid and a Terpenoid Hydrocarbon.—J. Piccard.—Cantharic acid is a powerful mono-basic acid. The author describes the copper and potassium salts, and the origin of cantharic acid, which he considers as formed from cantharic acid by the abscission of carbonic acid.

Gelatinous Silicic Acid and an Inorganic Membrane.—F. Ullik.—The author has found it possible to wash the well-known jelly produced by pouring dilute solution of soluble glass into hydrochloric acid and allowing the mixture to stand. He treats the mass repeatedly with water, taking care not to break it up, and has obtained it free from hydrochloric acid and chlorides, and in the formation of membrane-like layers, which, though incapable of filtering, are capable of dialysing like an animal membrane.

Contributions to the Voluminar and Steric Law.—H. Schröder.—Incapable of abstraction.

Action of Potassium Cyanate upon Epichlorhydrin.—A. L. Thomsen.—The reaction consists in the addition of 1 mol. cyanic acid to 1 mol. epichlorhydrin.

Bulletin de la Société Chimique de Paris,
No. 6, March 20, 1879.

Determination of the Total Nitrogen in Manures.—A. Rémont.—The author, referring to the well-known fact that when a manure contains nitrates the determination of its total nitrogen is impossible, recommends to add to the portion taken for analysis powdered white sugar to the extent of ten times the quantity of the nitrates supposed to be present, basing this approximation on the proportion of soluble matters which the manure contains. (If the manure contains soluble phosphates, salts of potassium, &c., the total soluble matter can throw no definite light on the quantity of nitrates which may be present.) The author adds the somewhat needless precaution that the sugared sample must be mixed with soda-lime before introduction into the combustion-tube lest detonation occur on heating.

Production of Crystalline Chromate of Baryta.—Leon Bourgeois.—Already noticed.

Certain Hydrocarbides formed by the Action of Methyl Chloride upon Toluene in Presence of Aluminic Chloride.—E. Ador and A. Rillet.—This memoir does not admit of useful abstraction.

Bromo-citraconic Acid.—Edme Bourgoin.—This acid, $C_{10}H_6Br_2O_3$, is bibasic, yielding alkaline and earthy alkaline salts of little stability and readily soluble in water.

Relative Affinities and the Reciprocal Displacements of Oxygen and the Halogens.—M. Berthelot.—The author proposes to show that the reciprocal displacements between the halogens and oxygen, united either with the metals or the non-metallic elements, may be predicted according to the sign of the quantities of heat liberated in the formation of the compounds which these elements form with oxygen on the one hand, and with the halogens on the other.

Thermo-chemical Researches on Magnesium, Calcium, Strontium, and Barium.—J. Thomsen.—Along with an increasing atomic weight we find an augmentation in—(a.) The stability of the hydrates and the heat liberated on the hydration of the oxides. (b.) The solubility of the hydrates in water and their heat of solution. (c.) The heat of formation of the chlorides, bromides, iodides, and nitrates. On the other hand, with an increasing atomic weight we find a decrease in—(a.) The affinity of the chlorides, bromides, iodides, nitrates, and sulphates for water; the proportion of water of crystallisation and the heat liberated on its fixation. (b.) The solubility and heat of solution of the same compounds. The atomic weight bears no relation to—(a.) The heat of neutralisation of the dissolved hydrates. (b.) The total heat of formation of the hydrates, i.e., R, O, H_2O .—*Journal f. Praktische Chemie*, xvi., p. 67.

Action of Mono-chloroacetic Acid upon the Sulphocyanates of the Aromatic Monamines.—J. H. Jæger.—Aniline and toluidine, dissolved in alcohol, have been treated with mono-chloroacetic acid along with ammoniac sulphocyanate.—*Fourn. f. Prakt. Chemie*, xvi., 17.

MISCELLANEOUS.

Mr. W. Valentin, F.C.S.—Mr. W. Valentin, F.C.S., is about to retire from the Laboratory of the Science Schools, South Kensington, formerly the College of Chemistry, in connection with which he has laboured for more than twenty years. It would be difficult to find one to whose patient work the present generation of chemists is more indebted, and a few of his old friends and pupils of the Royal School of Mines, recognising the value of his long services, are forming a committee with a view of presenting him with a testimonial. The Committee already includes Dr. Frankland, Dr. Forbes Watson, Prof. Guthrie, Dr. E. J. Mills, and Mr. W. Chandler-Roberts.

Iron and Steel Institute.—The Annual Meeting of the Iron and Steel Institute will be held on May 7, 8, and 9, at the Institution of the Civil Engineers. The programme includes papers and discussions on:—"The Mechanical Properties of Iron and Mild Steel," by Mr. D. Adamson, C.E. "On the Use of Steel in Naval Construction," by Mr. Nathaniel Barnaby, C.B., H.M.'s Chief Constructor. "On the Use of Steel in the Construction of Bridges," by Mr. H. N. Maynard. "On the Elimination of Phosphorus in the Bessemer Converter," by Mr. Sidney G. Thomas, F.C.S., and Mr. Percy C. Gilchrist, A.R.S.M., F.C.S. "On the Removal of Phosphorus and Sulphur during the Bessemer and Siemens-Martin Processes of Steel Manufacture," by Mr. G. J. Snelus, F.C.S., &c. "On a New Volumetric Method of Determining Manganese in Manganiferous Iron Ores, Spiegeleisen, Steel, &c.," by Mr. John Pattinson, F.I.C., Newcastle-on-Tyne. "On a Ready Means of Moulding Lime, and Making Lime or Basic Bricks and Linings for Furnace Converters, &c.," by Mr. Edward Riley, F.C.S., F.I.C., &c. "On a Practical Combination of the Bessemer and Puddling Processes," by Mr. Edwin Pettitt, Cheltenham. "On the Results of Working the Godfrey-Howson Furnaces at the Works of Tamaris, Gard, France," by M. Escalle. "On the Chemistry of Puddling," by Mr. H. Louis, A.R.S.M., Londonderry, Nova Scotia. "On a New Process for Protecting Iron and Steel against Rust," by Prof. Barff.

NOTES AND QUERIES.

Insurance of Tar Distilleries (Reply to W. A. R.).—Though not able to speak positively of the present time, I can certainly aver that a few years ago it was quite impossible to find any fire insurance office willing to grant the protection desired. I think it highly probable that the same state of things still exists, and, what is more, is extremely likely to do so. The tar distiller's best plan is to see to it that he sets such of his plant and materials requiring protection from the weather, under cover of the simplest and least inflammable kind. The tar-stills and other large stills worked with fire usually remain uncovered with any roof. My own experience is that with careful arrangement of plant, and proper precautions and care, there is little need for fear, and, in case of a fire even, this may, except in very exceptional cases, be quickly subdued without much damage being done. See communication on "Extinguishing Fires in Tar Distilleries, &c."—WATSON SMITH.

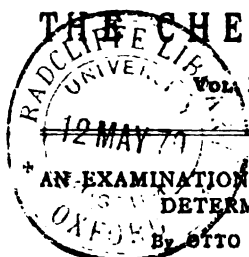
MEETINGS FOR THE WEEK.

MONDAY, 5th.—Medical, 8.30. (Anniversary)
— Royal Institution, 5. General Monthly Meeting.
— Society of Arts, 8. "Recent Advances in Telegraphy," by W. H. Preece. (Cantor Lectures)
TUESDAY, 6th.—Civil Engineers, 8.
— Royal Institution, 3. "Schumann," by Mr. Ernst Pauer.
— Zoological, 8.30.
WEDNESDAY, 7th.—Society of Arts, 8. "The Government Patent Bill," by W. Lloyd Wise, A.I.C.E.
THURSDAY, 8th.—Royal, 8.30.
— Royal Institution, 3. "Dissociation," by Prof. Dewar.
— Royal Society Club, 6.30.
— Society of Arts, 8. "The History of Alizarin and Allied Colouring Matters, and their Production from Coal-tar," by W. H. Perkin, F.R.S.
FRIDAY, 9th.—Royal Institution, 9. "Habits of Ants," by Sir John Lubbock.
— Astronomical, 9.
— Quekett, 8.
SATURDAY, 10th.—Royal Institution, 3. "Architecture," by Mr. H. H. Statham.
— Physical, 3.

TO CORRESPONDENTS.

E. W.—At the Library of the Commissioners of Patents, Southampton Buildings, Chancery Lane, W.C.
E. W. Young.—Such works as "Watts's Dictionary of Chemistry," "Ure's Dictionary of Arts, &c.," "Gannet's Physics," Wagner's and Richardson and Watts's works on Chemical Technology seem to already supply the information.

THE CHEMICAL NEWS.



VOL. XXXIX. No. 1015.

AN EXAMINATION OF DR. PAVY'S METHOD OF DETERMINING GLUCOSE.

By OTTO HEHNER, F.C.S., F.I.C.

THE glowing account given by Dr. Pavy (CHEMICAL NEWS, vol. xxxix., p. 77) of the results obtained in determining glucose by means of ammoniated Fehling solution has doubtless induced many chemists to repeat Dr. Pavy's experiments and to test the accuracy of his method.

120 c.c. of ordinary Fehling solution are mixed with 300 c.c. of strong ammonia (sp. gr. 0.88) and diluted to 1000 c.c. 100 c.c. of this liquid, containing 12 c.c. Fehling solution and capable of oxidising, according to Dr. Pavy, 0.05 grm. glucose, were used in the following experiments instead of 40 c.c., as recommended by Dr. Pavy, in order to obtain results comparable without calculation with those furnished by 10 c.c. of ordinary Fehling solution. The mode of titration was precisely that described by Dr. Pavy.

Following thus Dr. Pavy's directions I was surprised to obtain results widely differing from his, inasmuch as in a long series of experiments I found a difference of about 8 per cent in the total amount of glucose when determining it both by the ordinary copper solution and by the same rendered ammoniacal, the latter furnishing the higher results. In order to obtain corresponding results I had to dilute about 130 c.c. of Fehling solution with ammonia and water to 1000 c.c., instead of 120 c.c. as found by Dr. Pavy.

The copper solution used in these experiments contained 400 c.c. of soda solution (of 1.14 sp. gr.) per litre, and the sugar solution (inverted sugar) was quite neutral; hence there could not possibly be too large a quantity of alkali, against which Dr. Pavy warned in his paper.

I then prepared another quantity of Fehling solution, employing 480 c.c. soda, as recommended by Fresenius. I tested this both without and with the addition of ammonia, by titration with an invert sugar solution, obtained by heating 10153 grms. pure cane-sugar dissolved in 150 c.c. of water, with 10 c.c. strong HCl, on the water-bath, neutralising, and diluting to 250 c.c.

10 c.c. of Fehling solution required 11.70 c.c. sugar solution.

10 c.c. Fehling + 1 grm. NaHO used 11.65 c.c. sugar solution.

"Pavy Solution."	Sugar Solution.
100 c.c. used	11.22 c.c.
" + 0.2 grm. NaHO	11.40 "
" 0.4 "	11.55 "
" 0.6 "	11.71 "
" 0.8 "	11.70 "
" 1.0 "	11.72 "
" 2.0 "	11.90 "
" 3.0 "	12.08 "

The results show quite conclusively that Fehling solution prepared with 480 c.c. of soda of 1.14 sp. gr., corresponding to about 68 grms. NaHO per litre, does not give results concordant with the ratio stated by Dr. Pavy, and also that additions of very small quantities of soda steadily but perceptibly alter that ratio, until 50 grms. of soda per litre have been added to that already contained in the liquid, when further addition of soda, up to more than 80 grms., is without influence. Between these limits, namely with a total of 120 to 150 grms. of soda

per litre, Dr. Pavy's proportion holds strictly good, and then further addition of alkali again increases the oxidising power of the copper solution, but in much smaller proportion than before.

I further gradually diminished the alkali in the copper solution by adding measured quantities of standard acid, the results being as follow:—

Pavy Solution.	Sugar Solution.
100 c.c. — 0.20 NaHO required	11.00 c.c.
" — 0.40 "	10.71 "
" — 0.60 "	10.42 "
" — 0.81 "	8.80 "

Thus it appears that the smaller the quantity of fixed alkali, the smaller also the amount of sugar the copper solution is capable of oxidising; in fact, when no caustic soda at all is present, as in the last of the above experiments, 1 molecule of glucose requires exactly 8 molecules of cupric oxide for oxidation. With ordinary Fehling solution the ratio is 5 molecules; with the same rendered ammoniacal and containing no less than 120 grms. NaHO per litre, 6 molecules; and, lastly, the same without soda, 8 molecules. But the reaction without fixed alkali takes place exceedingly slowly, and on that account the end-point is not easy to hit off.

Instead, therefore, of guarding against too large a quantity of alkali, it is of far greater importance to avoid a deficiency, and good results can only be obtained by doubling the quantity of soda usually taken in the preparation of the cupro-tartrate solution. On the other hand, a much smaller excess of soda influences the result than is stated by Dr. Pavy, who found an addition of 5 grms. to 100 c.c. of the ammoniacal solution, or 417 grms. to the litre of Fehling solution without influence, whilst my results, which I have verified over and over again, show that even 2 grms. to 100 c.c. of the ammoniated solution, or 166 to the litre of Fehling, show a small but quite perceptible alteration. Thus the limits of alkali allowable or necessary are not so wide as might appear.

If the proportions given by me are adhered to, the method is capable of furnishing excellent results. The solution containing the glucose should be added drop by drop to the ammoniacal copper solution, kept gently simmering. The end-point of the reaction is beautifully sharp, and titrations of the same liquid will never differ by more than 0.1 c.c. of the sugar solution. The action is, however, somewhat slower than with ordinary Fehling liquid. By suitable arrangements the ammonia can readily be absorbed and any nuisance avoided which would result from its entering the atmosphere.

It must not, however, be taken for granted that the ratio holding good between the ordinary and the ammoniated copper solution is valid also in the case of other reducing saccharine matters. Dextrose and lævulose act equally, but I find that both milk sugar and maltose possess entirely different reducing power against two kinds of copper solutions.

Thus a solution of pure milk-sugar acted as follows:—

10 c.c. ordinary Fehling solution required	16.76 c.c.
100 c.c. Pavy solution	20.76 "

The ratio is therefore just contrary to that resulting by the action of glucose. The titration of milk-sugar, difficult and unsatisfactory as it is with the ordinary Fehling test, is, however, still more readily influenced by dilution, quantity of alkali, and rate of addition, and hence still less trustworthy in ammoniacal solution. Its action on that solution is very slow, and the end-point is somewhat difficult to observe, because on the experiment being much protracted cuprous oxide deposits and obscures the colour of the liquid.

Maltwort acts similarly to milk-sugar, proving that it does not contain glucose, but another sugar, doubtless maltose. Experiments in reference to this matter are in progress.

54, Holborn Viaduct, London, April 12.

RECENT CONTRIBUTIONS TO THE HISTORY
OF DETONATING AGENTS.*

By Professor ABEL, C.B. F.R.S.

(Continued from p. 188.)

It will now be evident why the readily yielding nature of the particles of liquid nitro-glycerine tends to counteract its great sensitiveness to detonation, and why, when the motion of the liquid particles is impeded by their admixture with solid matter, and when they are consequently placed in a position to resist mechanical motion by the force applied through the agency of detonation, its natural sensitiveness to detonation, and the rapidity with which it can be transmitted from particle to particle became fully developed.

Again, the reduction of gun-cotton fibre to a fine state of division, which renders the material readily convertible into very compact and dense masses, places the particles in the condition most favourable to resist mechanical motion upon the application of a blow, or of the concussion resulting from a detonation; hence, compressed gun-cotton is readily susceptible of detonation in proportion to the extent of compression, or to its density and compactness, while loose gun-cotton wool, or the lightly twisted or compressed material, cannot be readily detonated, because the force applied is expended in imparting motion to the readily yielding particles of the mass. If the force applied through the agency of a detonator to a mass of explosive material just borders upon that required for the development of the detonation, or if the condition of the mass is such as hardly to present the requisite resistance to mechanical motion essential for its detonation, then results intermediate between the mechanical dispersion of the mass and its violent chemical dispersion or disintegration, *i.e.*, detonation, are obtained. Thus frequent instances have been observed, especially in the experiments in the transmission of detonation through tubes, in which the initiative detonation has brought about an explosion attended with little, if any, destructive effect, portions of the mass being at the same time dispersed and occasionally inflamed. Not only have such results often been obtained with gun-cotton and dynamite, but even mercuric fulminate, exposed to the concussion of a distant detonation transmitted through a tube, has frequently been exploded in a manner quite distinct from the violent detonation developed in other instances. Silver fulminate, which under conditions detonates violently, even when only a particle of the mass is subjected to a sufficient disturbing influence, has been exploded without the usual demonstrations of force, by the transmitted effect of a detonation of mercuric fulminate. In these instances the violence of the concussion produced by the initiative detonation was only just bordering on that required for the development of detonation, and it appears probable that only some small portion of the mass operated upon was in a condition or position favourable to the action of the initiative blow. The remainder of the mass would then be dispersed by the gases developed from the detonated portion; in some instances the particles would be inflamed at the moment of their dispersion, in others they would even escape ignition. The latter appears to be always the case when gun-cotton is exploded by a blow from a hammer or falling weight. However carefully the arrangements are adjusted with a view to distribute such a blow uniformly over the entire mass struck, the concentration of a preponderance of the force applied upon some portion or portions of the entire mass, appears almost inevitable; hence only a small portion is actually detonated, the remainder being instantaneously dispersed by the gases suddenly generated while the weight is still resting upon the support.

Some experiments made in firing at masses of compressed gun-cotton, differently arranged and of different

thicknesses, with a Martini-Henry rifle, at short ranges, afforded interesting confirmation of the correctness of the explanation given of the operation of a blow upon masses of explosive material under different conditions. Disks of gun-cotton of the same density and diameter, but differing in thickness, were fired at; they were freely suspended, and their distance from the marksman was in all instances 100 yards. The thinnest disks were simply perforated by the bullets, not a particle of the gun-cotton being ignited. Somewhat thicker disks were inflamed by the impact of the bullet, while still thicker disks, fired at under the same conditions, were exploded, portions being in some instances dispersed in a burning state. No instance of detonation was, however, obtained. These differences in effect, obtained with masses of different thickness and weight, are due to the difference in their power to resist mechanical motion when struck by the bullet, and in the different amount of resistance to penetration presented by the thin and the thicker disks.

It has been explained that nitro-glycerin may be largely diluted with inert solid matters without the sensitiveness to detonation being reduced, while its detonation in open air becomes very much facilitated, because the mobility of the particles, and their consequent tendency to yield to the force of a blow or detonation, is very greatly diminished. But if a *solid* explosive agent is diluted with inert *solid* matter the case is different; for in such a mixture of the finely divided solid with non-explosive solid particles, there must be a partial and sometimes a complete separation of the particles of the explosive by the interposed inert particles with which it is diluted; hence the sensitiveness to detonation is reduced, and its transmission by the particles is retarded or altogether impeded, by a diminution of the extent of contact between the substance to be detonated and the initiative detonation, and by the barrier which the interposed non-explosive particles oppose to the transmission of detonation. Thus a mixture of mercuric fulminate with more than one-fifth its weight of French chalk could not be detonated by means of one grain of pure fulminate enclosed in a copper capsule, which was inserted into the mixture; that quantity, similarly confined, sufficed to detonate undiluted fulminate through a tube 8 inches long and 0.5 inch in diameter. In experiments made in this direction with finely divided gun-cotton, it was found that although dilution was an inert solid, applied in the *solid form*, reduced the sensitiveness of the material to detonation, this was not the case when it was incorporated with a salt soluble in water, the mixture being then compressed while in the wet state. The compressed masses thus obtained were, when dried, in a condition of greater rigidity than could be attained by submitting undiluted gun-cotton to considerably more powerful pressure, because the crystallisation of the soluble salt used as the diluent upon evaporation of the particles, cemented the particles composing the mass more rigidly together. The gun-cotton was therefore presented in a form more capable of resisting the mechanical action of a small charge of fulminate than a more highly compressed undiluted gun-cotton, and hence the reduction in sensitiveness due to the detonation of the explosive compound is nearly counterbalanced by the greater rigidity imparted to the mass. If a soluble oxidising agent (a nitrate or chlorate) be employed as the diluting material, the predisposition to chemical reaction between it and the gun-cotton (which is susceptible of some additional oxidation), appears to operate in conjunction with the effect of the salt in imparting rigidity to the mixture, thus rendering the latter quite as sensitive to the detonating action of the minimum fulminate charge as undiluted gun-cotton. Moreover, the interesting fact has been conclusively established that these compressed mixtures of gun-cotton with a nitrate or a chlorate are much less indifferent to the influence of detonating nitro-glycerin than gun-cotton in its pure state. Chlorated and nitrated gun-cotton are detonated with certainty by means of $\frac{1}{2}$ oz. of nitro-glycerin, whereas the detonation of 2 ozs. of the latter

* Abstract of a Paper read before the Royal Institution of Great Britain, Friday, March 27, 1879.

accomplished the detonation of ordinary compressed gun-cotton only once in a large number of experiments.

If compressed gun-cotton is diluted by impregnating the mass with a liquid, or with a solid which is introduced into the mass in a fused state, its susceptibility of detonation is reduced to a very much greater extent than by a corresponding quantity of a solid inert body, incorporated as such with the gun-cotton, the cause being the converse of that which operates in preventing a reduction of the sensitiveness to detonation of nitro-glycerin by its dilution with an inert solid. In this case the explosive liquid envelopes the solid diluent, and remains continuous throughout, occupying the spaces which exist between the solid particles; hence detonation is readily established and transmitted. But in the case of the solid explosive, the diluent, which is liquid, or at any rate is introduced into the mass in the liquid state, envelopes each particle of the solid, so that a film of inert material surrounds each, isolating it from its neighbours, and thus opposing resistance to the transmission of detonation, which is proportionate to the original porosity or absorbent power of the mass.

While compressed gun-cotton, in the air-dry state, is detonated by 2 grains of mercuric fulminate imbedded in the material, its detonation by 15 grains, applied in the same manner, becomes doubtful when it contains 3 per cent of water, over and above the 2 per cent which exists normally in the air-dry substance. Specimens which had been impregnated with oil or soaked in melted fat and allowed to cool could not be detonated by means of 15 grains of fulminate. These diluted samples of gun-cotton could only be detonated by adding very considerably to the power of the initiative detonation; 100 grains of confined fulminate generally failed to detonate gun-cotton containing from 10 to 12 per cent of water, and if the amount reached 17 per cent, 200 grains of fulminate were needed to ensure its detonation.

But moist or wet compressed gun-cotton is decidedly more susceptible of detonation by (dry) compressed gun-cotton itself than by mercuric fulminate.

Thus 100 grains of dry gun-cotton, detonated through the agency of the ordinary fulminate fuze, suffice to detonate wet gun-cotton containing 17 per cent of water, though this result is somewhat uncertain. If the diluting agent amounts to 20 per cent, detonation is not certain with less than 1 oz. of dry gun-cotton, and if the compressed material be completely saturated with water (*i.e.*, containing 30 to 35 per cent) 4 ozs. of the air-dry substance, applied in close contact, are needed to ensure its detonation.

Detonation is transmitted through tubes from dry compressed gun-cotton to a moist disk of the material with the same facility as to the dry substance; and this is also the case with regard to the propagation of detonation from one mass of moist gun-cotton to another, in open air, all the pieces being ranged in a row, in contact with each other, provided that the piece first detonated does not contain less water than the others to which detonation is transmitted. Some curious results, obtained in experiments on the transmission of detonation, with gun-cotton containing different proportions of water, appeared to indicate that the character or quality of detonation developed by gun-cotton is subject to modification by the proportion of water which the latter contains.

Gun-cotton containing 12 to 14 per cent of water is ignited with much difficulty on applying a highly heated body. As it leaves the hydraulic press upon being converted from the pulped state to masses having about the density of water, it contains about 15 per cent of water; in this condition it may be thrown on to a fire or held in a flame without exhibiting any tendency to burn; the masses may be perforated by means of a red-hot iron or with a drilling tool, and they may with perfect safety be cut into slices by means of saws revolving with great rapidity. If placed upon a fire and allowed to remain there, a feeble and transparent flame flickers over the surface of the wet

gun-cotton from time to time as the exterior becomes sufficiently dry to inflame; and in this way a piece of compressed gun-cotton will burn away very gradually indeed. A pile of boxes containing in all 6 cwts. of gun-cotton, impregnated with about 20 per cent of water, when surrounded by burning wood and shavings in a wooden building, was very gradually consumed, the gun-cotton burning as already described when the surfaces of the masses became partially dried. In two other experiments quantities of wet gun-cotton of 20 cwts. each, packed in one instance in a large, strong wooden case, and, in the other, in a number of strong packing cases, were placed in small magazines, very substantially constructed of concrete and brickwork. Large fires were kindled around the packages in each building, the doors being just left ajar. The entire contents of both buildings had burned away, without anything approaching explosive action, in less than two hours. This comparatively great safety of wet gun-cotton, coupled with the fact that its detonation in that condition may be readily accomplished through the agency of a small quantity of dry gun-cotton, which, through the medium of a fulminate fuze or detonator, is made to act as the initiative detonating agent, gives to gun-cotton important advantages over other violent explosive agents for purposes which involve the employment of more or less considerable quantities at one time, on account of the comparative safety attending its storage and the necessary manipulation of it. Moreover, it has been well established by experiments of many kinds carried out on a considerable scale, as well as by accurate scientific observations, that the detonation of wet gun-cotton is decidedly sharper or more violent than that of the dry material; a circumstance which affords an interesting illustration of the influence exerted by the physical condition of the mass upon the facility with which detonation is transmitted from particle to particle. In the determinations made by means of the Noble chronoscope, of the velocity with which detonation is transmitted along layers or trains of gun-cotton and nitro-glycerin, the lecturer has included experiments with gun-cotton containing different proportions of water. When the material contained 15 per cent of the liquid, some indications were obtained that the rate of transmission of detonation was a little higher than with dry gun-cotton; the difference was very decidedly in favour of wet gun-cotton, when the latter was thoroughly saturated with water. (With air-dry gun-cotton the mean rate of transmission ranged in several experiments between 17,000 and 18,900 feet per second; with gun-cotton containing about 30 per cent of water, the mean rate of transmission ranged between 19,300 and 19,950 feet per second.) The air in the masses of compressed gun-cotton being replaced entirely by the comparatively incompressible body, water, the particles of explosive are in a much more favourable condition to resist displacement by the force of the detonation, and hence they are more readily susceptible of sudden chemical disintegration. Moreover, the variations in the rate of travel of detonation in dry gun-cotton, resulting from differences in the compactness or rigidity of different masses of the material, are very greatly reduced, if not entirely eliminated, by saturating the disks with water and thus equalising their power of resisting motion by a sudden blow.

Another striking illustration of the influence which the physical character of an explosive substance exercises over its susceptibility to detonation and the degree of facility with which its full explosive force is developed, is furnished by one of the most recently devised, and one of the most interesting of existing, explosive agents.

Twelve years ago, soon after the process of producing compressed and granulated gun-cotton had been elaborated by the lecturer, it occurred to him to employ these forms of gun-cotton as vehicles for the application of nitro-glycerin. A considerable proportion of the liquid was absorbed by the porous masses of gun-cotton, and a nitro-glycerin preparation analogous in character to dynamite was

thus obtained. The absorbent was in this case a violently explosive body instead of an inert solid as in dynamite, but the quantity of nitro-glycerin in a given weight of the preparation (to which the name of *Glyoxilin* was given), was considerably less than in the Kieselguhr-preparation; hence the latter was nearly on a point of equality with it, in regard to power, as an explosive agent.

Nobel has observed that if, instead of making use of the most explosive form of gun-cotton, or trinitro-cellulose, a lower product of nitration of cellulose (the so-called soluble or colloid gun-cotton) is added to nitro-glycerin, the liquid exerts a peculiar solvent action upon it, the fibrous material becoming gelatinised while the nitro-glycerin becomes at the same time fixed, the two substances furnishing a product having almost the characters of a compound. By macerating only from 7 to 10 per cent of soluble gun-cotton with 90 to 93 per cent of nitro-glycerin, the whole becomes converted into an adhesive plastic material, more gummy than gelatinous in character, from which, if it be prepared with sufficient care, no nitro-glycerin will separate even by its exposure to heat in contact with bibulous paper, or by its prolonged immersion in water, the components being not easily susceptible of separation even through the agency of a solvent of both. As the nitro-glycerin is only diluted with a small proportion of a solidifying agent which is itself an explosive (though a somewhat feeble one), this *blasting gelatine*, as Nobel has called it, is more powerful not only than dynamite but also than the mixture of a smaller quantity of nitro-glycerin with the most explosive gun-cotton, as the liquid substance is decidedly the most violent explosive of the two. Moreover, as nitro-glycerin contains a small amount of oxygen in excess of that required for the perfect oxidation of its carbon and hydrogen constituents, while the soluble gun-cotton is deficient in the requisite oxygen for its complete transformation into thoroughly oxidised products, the result of an incorporation of the latter in small proportion with nitro-glycerin is the production of an explosive agent which contains the proportion of oxygen requisite for development of the maximum of chemical energy by the complete burning of the carbon and hydrogen, and hence this *blasting gelatin* should, theoretically, be even slightly more powerful as an explosive agent than pure nitro-glycerin.

That such is the case has been well established by numerous experiments, but although this *blasting gelatin* may be detonated like dynamite by means of small quantities of confined detonating composition, when it is employed in strongly tamped blast-holes, or under conditions very favourable to the development of great initial pressure, it behaves very differently from that material, or other solid though plastic preparations of nitro-glycerin, if the attempt is made to detonate it when freely exposed to the air or only partially confined. It not only needs a much more considerable amount of strongly confined detonating composition than dynamite and similar preparations do, to bring about a detonation with it under those conditions; but when as much as 15 or 20 grains of confined fulminate are detonated in direct contact with it, although a sharp explosion occurs, little or no destructive action results, and a considerable portion of the charge operated upon is dispersed in a finely-divided condition. This dispersion appears to take place to some slight extent with dynamite also, when a small charge is detonated in open air, in consequence of its want of rigidity, though the amount of explosive which thus escapes detonation is very small as compared with the gelatin.

(To be continued.)

Temper of Steel.—M. Jarolimek maintains that steel can not only be hardened by immersion in boiling water, but even in boiling oil, melting lead, and melting zinc.—*Moniteur Scientifique.*

EXHIBITION OF ELECTRIC LIGHTING APPARATUS AT THE ROYAL ALBERT HALL.

ON Wednesday evening last an Exhibition of Electric Lighting Apparatus at the Royal Albert Hall was inaugurated by a lecture on the electric light, which was given by Mr. W. H. Preece, M.Inst.C.E., in the presence of the Prince of Wales, the Duke of Edinburgh, Prince Christian, and the Duke of Teck. Among the visitors were Lord Lindsay, Lord Rayleigh, Sir F. Pollock, Sir Charles Bright, Dr. Lyon Playfair, C.B., F.R.S., Dr. De la Rue, F.R.S., Prof. Tyndall, F.R.S., Prof. Abel, C.B., F.R.S., Dr. William Siemens, F.R.S., Dr. Frankland, F.R.S., Mr. Crookes, F.R.S., Mr. J. N. Lockyer, F.R.S., Prof. Ayrton, &c.

After the Prince of Wales had taken a seat on the right of the orchestra, the Duke of Edinburgh advanced to the front of the platform, and as Senior Vice-President of the Council of the Albert Hall introduced the Lecturer, who began by demonstrating that all systems of artificial illumination are dependent upon the production of heat. Some brilliant experiments were introduced to show that the greatest concentration of heat can be produced by electric currents in overcoming resistance. When this resistance is the air itself the electric arc is the result: when it is a wire or carbon rod we have light by incandescence. The discussion at the present time was whether light by the arc or by incandescence is to be the light of the future. The arc gives two and a half times the intensity of light given by incandescence by the same power. Mr. Preece showed that the improved methods now employed for the production of electricity by mechanical means is due to the development of Faraday's discovery that electric currents are produced by the rapid rotation of coils of wire in a powerful magnetic field, Faraday's original apparatus for showing this being exhibited by the Royal Institution. The electric light dependent on incandescence was practically illustrated by the lamps of Werdermann and the Anglo-American Light Company. That dependent on the arc was shown by the regulators of Serrin, Siemens, Rapieff, and Wallace. The form of light based on the arc, but independent of regulators, called the candle, was illustrated by those of Wilde and Jablochhoff. Mr. Preece also introduced to the audience a powerful lamp called the "holophote," which is about to be introduced into the ports at Spithead, with a view to testing their value in detecting the advance of an enemy's torpedo. The many shortcomings of the electric light—such as the noise, the flickering, the deep shadows—were referred to, as well as such advantages as the absence of smoke, and the purification instead of the poisoning of the air in large buildings. The electric light was pronounced to be at the present time in the tentative stage, there being, as Mr. Preece remarked, three stages of every physical invention—the theoretic, when it is the dream of the philosopher; the tentative stage, when it is the dream of the capitalist; and the third stage, when the practical man realises these dreams.

The lecture was very warmly applauded. At its conclusion the Duke of Edinburgh proposed, on behalf of the Council of the Royal Albert Hall, a vote of thanks to Mr. Preece for his lecture. In the course of his speech His Royal Highness said that the Council of the Royal Albert Hall was anxious to employ the electric light even in its present imperfect condition in lighting the building, and would persevere until something satisfactory was accomplished. The vote of thanks was seconded by His Royal Highness the Prince of Wales.

The Exhibition will remain open until Saturday, and on Friday evening, at 8 o'clock, an explanatory lecture will be given by Mr. J. N. Shoolbred. The Council has endeavoured to make the collection of apparatus embrace the entire scope of all applications into which electric light strictly enters; and also as completely represents

tive as possible of all the various methods hitherto devised for the production of the electric light by mechanical means.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, May 1, 1879.

Dr. WARREN DE LA RUE, President, in the Chair.

THE minutes of the previous meeting were read and confirmed. The following certificate was read for the first time:—J. Sakurai.

THE PRESIDENT then called on Dr. W. RAMSAY to read a paper "*On the Volumes of Liquids at their Boiling-points obtainable from Unit Volumes of their Gases.*" Kopp, in 1855, pointed out that the specific gravities of organic compounds show a certain regularity with regard to each other. If the molecular weights of various compounds be divided by their respective specific gravities at their boiling-points, a series of numbers is obtained, which Kopp ultimately named specific volumes. Kopp's method of determining the volume of a liquid at its boiling-point (the only point at which volumes are comparable, for at that point the vapour-tensions of all liquids are equal) was to ascertain the boiling-point with great accuracy, to determine the specific gravity of some known temperature, and calculate the volume required by means of the coefficient of expansion. This process involves the use of complicated and expensive apparatus, and necessitates laborious calculation. Before describing the apparatus used by himself, the author discusses the precise conceptions involved in the expressions used by Kopp, &c.—"Atomic volume," "molecular volume," and "specific volume:" specific volume as used by Kopp =

$$\frac{\text{molecular weight}}{\text{specific gravity}}$$

By molecular weight is meant the specific gravity of the gas, hydrogen at 0° being equal to 1; specific gravity water = 1; so that two scales are employed, viz., one based on H₂ as unity, and one based on water. The number obtained from the above equation, therefore, does not give the relation between the volumes and weights of liquids at their boiling-points and that of the gases obtainable from those liquids. To obtain this relation, the number representing the specific gravity of the liquid must be multiplied by the number which represents the relation between the specific gravity of water and that of hydrogen, i.e., 22.326. By reversing Kopp's process, i.e., by dividing the sp. gr. of the boiling liquid by that of the gas, the amount of gas obtainable from unit volume of the liquid is calculated. Thus, specific volume of liquid hydrogen

= 5.5, and its sp. gr. is $\frac{1}{5.5}$ or 0.1818, and $0.1818 \times 22.326 = 4.0592$, the sp. gr. of liquid hydrogen (hydrogen gas = 1). By dividing the sp. gr. of gaseous hydrogen by that

of liquid hydrogen, or $\frac{1}{5.0592}$ we get 0.0024635 as the volume of liquid obtainable from 1 volume of gas, or 2.46 from 10,000 vols. of gas. This number (2.46) the author proposes to call "ebullition volume." Kopp's numbers give the amounts of boiling liquids obtainable from 22.326 vols. of gas at 0°. The apparatus used by the author consists of a thin glass bulb of about 10 c.c. capacity, in shape like a lemon: its upper end is terminated by a hooked capillary tube, and its lower end is closed, and furnished with a glass hook. The capacity is accurately determined by filling with boiled distilled water and weighing. To determine the weight of a known volume of a liquid at its

own boiling-point, the bulb is filled with the liquid, and suspended by a platinum wire in a glass vessel resembling a large test-tube, having a bulb blown about half an inch from the bottom. A small quantity of the liquid is placed in this glass flask, and the bulb suspended by a platinum wire from the cork which fits in the upper part of the flask. The liquid in the flask is caused to boil by a small Bunsen flame: the vapour heats the bulb, the liquid in it expands, and drops are expelled from the capillary tube. As soon as the drops cease to fall, usually in about ten minutes, the bulb and its contents have assumed a temperature equal to the boiling-point of the liquid. The bulb is withdrawn, allowed to cool, and weighed. Various precautions must be taken with liquids which are very volatile or oxidisable. Allowance is carefully made for the expansion of the glass. The time required for one determination is about half an hour. The author has made many determinations with a great variety of substances. His results, on the whole, agree very closely with those obtained by Kopp. The value of some of the elements seems to vary much. Thus, oxygen has two values, 3.49 and 5.45; sulphur 10.27 and 12.79. Nitrogen in amines is 2.3; in cyanogen compounds, 1.7; in bodies containing NO₂, 1.74 (Kopp). The author has determined the ebullition volume of some of the pyridine series of bases, e.g., in picoline, value of N = 4.08; in its isomeride aniline, N = 2.11; in pyrrol N = 9.12. From its volume it appears to belong to the cyanogen group. In conclusion, the author draws attention to the enormous differences between these values of N, and suggests a connection between values and the amounts of heat evolved in the formation of these substances.

The next paper was read by Mr. J. PATTINSON, "*On a Method of Precipitating Manganese entirely as Dioxide, and its Application to the Volumetric Determination of Manganese.*" Many methods of determining manganese volumetrically have been suggested, but none have come into general use, owing principally to the difficulty of obtaining the whole of the manganese in a definite and uniform state of oxidation. The author has examined the methods suggested by Pereno and Lenssen, but did not succeed in obtaining regular results. Wright and Luff have also been unable to obtain pure manganese dioxide by any of the ordinary methods. After numerous experiments the author found that the whole of the manganese in a solution of manganous chloride could invariably be precipitated in the condition of dioxide, if a certain amount of ferric chloride be present, by a sufficient excess of a solution of calcium hypochlorite or bromine water, adding, after heating the solution to from 140° to 160° F., an excess of calcium carbonate, and then well stirring the mixture. Without the ferric salt the precipitation as MnO₂ is imperfect. Zinc chloride may be substituted for ferric chloride, but neither aluminium nor barium chlorides have the same desirable effect. The author recommends the following solutions, &c.:—The clear liquid obtained by decantation from a 1.5 per cent solution of bleaching-powder; light granular calcium carbonate obtained by precipitating an excess of calcium chloride by sodium carbonate at 180° F.; a 1 per cent solution of ferrous sulphate in dilute (1 in 4) sulphuric acid; standard solution of potassium dichromate equivalent to 1 part of iron in 100 of solution. The application of the process to manganiferous iron ores is as follows:—10 grains of the ore, dried at 212°, are dissolved in a 20-oz. beaker in about 100 fluid grains of hydrochloric acid (sp. gr., 1.18). Calcium carbonate is then added until free acid is neutralised and the liquid turns slightly reddish. 6 or 7 drops of HCl are now added, and 1000 grains of the bleaching-powder solution, or 500 grains of saturated bromine-water, and boiling water run in until the temperature is raised to 140° to 160° F.; 25 grains of calcium carbonate are added, and the whole well stirred. If the supernatant solution has a pink colour, the permanganate is reduced by a few drops of alcohol. The precipitated oxides of iron and manganese are filtered off and washed. 1000 grains of the acidified ferrous sulphate

solution are carefully measured into the 20-oz. beaker already used; the filter with its washed contents added. A certain quantity of the ferrous sulphate is oxidised by the MnO_2 ; this quantity is estimated with the standard dichromate solution, when the quantity of MnO_2 can easily be calculated. The iron present must be at least equal in weight to the manganese during the precipitation in order to ensure the absence of lower oxides. The author gives in detail the slight modifications necessary in the analysis of spiegeleisen, ferro-manganese steel, and manganese slags; also some analyses which prove the accuracy of the process.

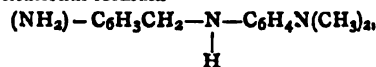
Dr. WRIGHT was extremely interested in the paper, especially as to the action of the ferric chloride. He asked if Mr. Pattinson had found any substances which acted in the opposite way, i.e., decreased the yield of MnO_2 .

Mr. PATTINSON said that all his efforts had been directed to ensure the complete formation of MnO_2 .

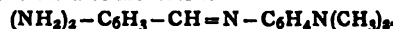
The PRESIDENT then called on Mr. WARINGTON to read a paper "On the Determination of Nitric Acid as Nitric Oxide by means of its Action on Mercury." During the last two years the author has used this method to a considerable extent: it was first suggested by W. Crum, and has been much improved by Frankland. The author has in the main carried out the process as recommended by Thorp in Sutton's "Volumetric Analysis." He did not, however, expel any gas which might be liberated before shaking. In the presence of some kinds of organic matter a permanent froth was at some times produced. This could be destroyed either by the introduction of a little hot water through the stopcock, or by gently warming the liquid in the inclined tube by a small flame. A considerable number of experiments are given as to the effect of chlorides on the accuracy of the results obtained in estimating nitrates, nitrites, and the nitric acid in soils. The author finds that the results are quite as satisfactory, even when chlorine is present in a quantity equal to eight times the equivalent of the nitrogen, and concludes that chlorides, except perhaps in extreme cases, are no hindrance to the accurate determination of nitric and nitrous acid by this method, and that it is unnecessary to remove them by previous treatment with sulphate of silver. The author then investigated the influence of organic matter, and finds that organic matter in quantities likely to be met with affect the results but little; cane-sugar, however, has a remarkable action in preventing the complete evolution of the nitric oxide. In some cases a considerable quantity of gas was evolved, without shaking; this gas was found to be nitric oxide, the action taking place between the nitric acid and organic matter instead of between nitric acid and mercury. It is obvious that a considerable error would be caused by expelling any gas liberated before shaking in such cases. Commercial glucose has a considerably less injurious effect than cane-sugar. It is interesting to observe that chlorides, if present, seem to prevent to a great extent the injurious effect of cane-sugar. Thus, a solution containing 20.8 parts of nitrogen gave by the process, after removing chlorides, 12.6 and 11.5, but with chlorides not removed, 18.5. In some cases the author has observed that the reaction is not completed in the shaking-tube, but that bubbles of gas continue to be evolved when the liquid was transferred to the measuring-tube; the reason of this defective reaction was not discovered. The author draws the following conclusions:—That in the absence of organic matter, and with proper manipulation, the method is one of great accuracy, and is capable of determining extremely small quantities of nitrates and nitrites; the natural error of the process is a small one of deficiency; the presence of chlorides in moderate quantity is no hindrance; quantities of organic matter, small in relation to the nitrates present, have little or no effect on the results; larger quantities, especially of cane-sugar, may cause a considerable deficiency; this deficiency is reduced by the presence of chlorides, but is not entirely removed.

After a few remarks by Dr. GILBERT,

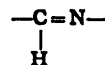
The PRESIDENT called on Dr. O. N. WITT to read a paper "On a New Class of Colouring Matters." This is the first part of a series on the simultaneous gradual oxidation of amido and methyl groups. If a mixture of meta-toluylen-diamin and dimethyl-para-phenylen-diamin be oxidised in aqueous solution, or if the formation of the latter of these bases from nitroso-dimethyl-aniline be combined with this oxidation process, by acting upon meta-tolulylen-diamin with nitroso-dimethyl-anilin, the reaction results in either case in the formation of a new compound of an intense blue colour, for which the author proposes the name of toluylen blue, $\text{C}_{15}\text{H}_{16}\text{N}_4\text{HCl} + \text{H}_2\text{O}$. It is the neutral salt of a new triatomic base, the acid salts being reddish brown. If treated with reducing agents it absorbs 2 atoms of hydrogen, forming a new colourless base, having the constitutional formula—



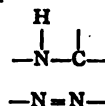
and the formula of the blue is—



This compound may be considered as the first representative of a new class of colouring matters, which have the group—



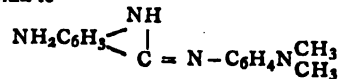
as a chromophor, and form, from their properties and constitution, a connecting link between the rosanilin and the azo series, the chromophor of the former being—



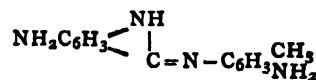
and of the latter—



One of the most remarkable properties of toluylen blue is its power of absorbing hydrogen from other amines, and being reduced to its leuko-compound, the amine undergoing condensation similar to that effected by oxidising agents and invariably resulting in the formation of new colours. Thus the blue, by acting on another molecule of itself produces toluylen pink, $\text{C}_{15}\text{H}_{16}\text{N}_4$, a crystallised compound forming two series of salts, one pink the other blue. By acting with toluylen blue on meta-tolulylen-diamin for twelve hours at 35° toluylen violet is obtained, $\text{C}_{14}\text{H}_{14}\text{N}_4$. This new compound dissolves sparingly in alcohol and ether with a pink colour, exhibiting a most brilliant fluorescence; it forms two series of salts, one violet the other green. The constitutional formula of toluylen pink is—



and of the violet—



The general conclusions which may be drawn are that under the above circumstances a gradual dehydrogenisation of the amido- and the methyl-groups takes place, the quantities of hydrogen removed from both sources being always the same; that in a great many cases the use of oxidising agents may be avoided by using the corresponding nitroso-compounds, or by utilising the tendency of some organic compounds for the absorption of hydrogen. The author promises further investigations in the same direction. Specimens of the above new and beautiful colouring-matters were exhibited by the author.

The Society then adjourned to May 15, when the following papers will be read:—"On Nitrification," Part II., by R. Warington; "On Alkaloids of the Veratrum,"

Part III., by Dr. Wright and Mr. Luff; "On Alkaloids of the Veratrums," Part IV., by Dr. Wright; "Alkaloids of the Aconites (Part IV.), and on Japanese Aconite Roots," by Dr. Wright and Mr. Luff; "On the Action of Hydrochloric Acid on Manganese Dioxide," by Spencer Pickering; "The Composition of Milk in Health and Disease," by A. Wynter Blyth; "Notes on the Effect of Alcohol on the Chemistry of Digestion," by W. H. Watson.

NOTICES OF BOOKS.

A Dictionary of Chemistry, and the Allied Branches of other Sciences. By HENRY WATTS, F.R.S., F.C.S., Editor of the *Journal of the Chemical Society*; assisted by Eminent Contributors. Third Supplement, Part I. Longmans and Co., 1879.

THIS is the third supplementary volume of "Watts's Dictionary" which has been issued since the completion of the body of that work, and brings the record of chemical discovery down to the end of the year 1877, including some of the more important discoveries which have appeared in 1878. The volume, which begins with "Absinthol" and ends with "Fustic," comprises only one-quarter of the letters of the alphabet; we do not see, therefore, how the other three-quarters can be compressed into the single volume mentioned in the Preface. It is true that the articles on those very fruitful bodies the benzo-compounds take up no less than one-fifth of the part, but, notwithstanding this, if the remainder of the alphabet is to be squeezed into a single volume, it will have to be a very bulky one, or else an injurious amount of compression will have to be exercised. The first important article we meet with is on Aniline Colours, which gives the best method of testing their tinctorial power by an accurately titrated solution of sodic hyposulphite. We also have accounts of the formation of aniline-black by the action of vanadic salts and by electrolysis; also of its composition and probable formula. The article also treats of aniline-green, grey, and red. Anthracen very naturally claims a large amount of attention, the newest views as to its molecular constitution being given at length. Its latest derivatives and its isomerides are then described, as well as its fluorescent properties. Under Anthraquinon we have a full account of the most recently discovered derivatives of that prolific body, and of the formation of artificial alizarin from pyrocatechin. Under this heading are also described quinizarin, purpurloxanthin, the anthroflavic acids, the anthroflavons, chrysazin, purpurin, anthrapurpurin, flavo-purpurin, oxy-chrysazin, and rubopin. Under Arsenic the continuation of Nilsson's classical researches on the compounds of that metal are described, more especially those on its sulphur salts. In the article Atmosphere we have the most recent determinations of the amount of carbon dioxide, ammonia, hydrogen peroxide, and dust in the air by various observers. A few pages further on we have an exhaustive article on barley, from Mr. R. Warington, the most recent results obtained by Messrs. Lawes and Gilbert being fully described. Under Beer we have full directions for the detection of foreign bitters in that beverage, from which we learn that brewers use nearly thirty different "hop-substitutes," more or less poisonous, from strychnine and brucine to gentian and wormwood. Benzacetic acid introduces us to the benzo-compounds, to which no less than 167 pages are devoted. It is strikingly indicative of the progress of research in this direction that in the original work the account of these compounds from their first discovery onwards only took up 40 pages, while in the first and second supplements we have 82 and 57 pages respectively given to these bodies. In these articles the vexed question of the determination of the relative posi-

tions of the substituted radicles in benzene-derivatives is fully discussed, the views of Körner, Graebe, and others being given. The constitution and structure of the various mono-, di-, tri-, and tetra-bromo-benzene, as well as of their nitro- and chloro-congeners, are also thoroughly investigated. This article likewise contains an account of the iodo-, bromo-, and chloro-derivatives of aniline, the amido-benzenes, and the azo-benzenes. A useful table is given of the abbreviated symbol, the physical properties and positions of the substituted radicles of more than 120 ortho-, meta-, and para-benzene derivatives. The next part of this article treats of benzene-sulphonic acid and its numerous derivatives, and gives a table of the physical properties of its principal ortho-, meta-, and para-salts. A number of other tables are also given of the chemical and physical properties of the derivatives of benzene-meta-disulphonic acid and of those of the brom-amido-benzene-disulphonic acid. Benzoic acid and its numerous derivatives are next treated of, the newest methods of making ortho-oxy-benzoic acid or salicylic acid being fully described. Directions are added for the purification, detection, and estimation of this now important compound. The next important articles are those on the inorganic and organic compounds of boron. Under Cæsium we have Godeffroy's method of separating cæsium, rubidium, and potassium by preparing their alums and separating them by fractional crystallisation according to the process originally devised by Redtenbacher. The Cerite group of metals also receives ample treatment, a large number of their principal salts being fully described. The article on Chemical Action is an important one from a philosophical point of view, the velocity of chemical action, the retardation of chemical reactions by indifferent substances, the decomposition of certain salts by water, the mutual replacement of halogen elements, and other similar questions being discussed at great length. Chloral and its compounds have 8 pages devoted to them, while cinchona barks, bases, and compounds take up 26 pages. Cinnamic and citric derivatives also receive their full share of attention. Thomas's researches on the gases enclosed in the coal of the South Wales basin are given nearly in full under Coal. Cumulative Resolution is a highly philosophical article from the pen of Dr. Mills. The other important articles are on Flame, by Professor Thorpe; Forest Trees, by Professor Warington; on the Diphenyl Derivatives, which have nearly 30 pages given to them; Dye Stuffs, Electricity, Electro-capillarity, Ethyl Derivatives, Fermentation, Fulminates, and Fulminurates.

For the second part, which we trust will shortly appear, we are promised contributions by Drs. Armstrong, Roscoe, Thorpe, Flight, and Professors G. C. Foster and R. Warington.

The Chemistry of Common Life. By the late JAMES F. W. JOHNSTON, Professor of Chemistry in the University of Durham, &c. A New Edition, revised and brought down to the present time, by ARTHUR HERBERT CHURCH, M.A. Oxon, &c. Blackwood and Sons, 1879.

JUST twenty years have elapsed since that versatile writer the late Mr. G. H. Lewes took up the pen which, so to speak, had fallen from Prof. Johnston's hand some two or three years before, and brought a second edition of a book which, amongst popular scientific works, had already taken its stand by the side of "Liebig's Letters on Chemistry." Prof. Johnston had performed his task so thoroughly that Mr. Lewes's utmost diligence had not been able to glean much after him, although seven years had passed between the appearance of the original and the revised editions. The book had three great merits—it was written in a popular and entertaining style, it was comprehensive in the choice of subjects, and, above all, it was thoroughly exact. It seems, therefore, strange that a couple of decades should have been allowed to pass

away between the publication of the second and third editions.

Mr. Church has executed his task as a pious duty, and has been most careful to respect the method, the style, and the matter of Prof. Johnston's original, only such corrections and alterations having been made as were rendered necessary by the recent advances made in scientific knowledge. Only very few additions have been made, one of them of notable value, in the form of a chapter on the Colours we Admire, which forms a fitting pendant to the well-known chapters on the Odours we Enjoy. A sketch is given of the origin and properties of hæmaglobin turacin, the cupreous pigment discovered by Mr. Church in the red colouring matter of the wing feathers of some eleven species of the Touraco, the black pigment found in black hair and feathers, chlorophyll, coelein, alizarin, and, lastly, of the coal-tar dyes.

In bringing out the third edition of this work Mr. Church possessed an advantage over the late Mr. Lewes, for he had the opportunity of consulting Prof. Johnston's private and corrected copy of the "Chemistry of Common Life," so that he was not only enabled to incorporate in his revisions some valuable matter gathered by the late author, but to learn the kind of additions which he contemplated.

Although appearing in one volume instead of two, as in the case of the first two editions, the present issue contains more matter, the type being smaller and closer and the pages larger.

OBITUARY.

WILLIAM GEORGE VALENTIN, F.C.S.

LAST week we stated that a committee had been formed for the purpose of presenting Mr. Valentin with a testimonial in recognition of his long services to chemical science. It is now our painful duty to announce that he died suddenly on the 1st instant from apoplexy. This well-known chemist was born at Neuenburg, in the Black Forest, on the 16th of May, 1829. He came to England in 1855, and in the early days of the College of Chemistry studied under Dr. Hofmann, who esteemed him greatly, and, recognising his ability, made him Senior Assistant in the Laboratory. He retained this position under Dr. Frankland, with whom he organised the new laboratories of the Science Schools, South Kensington. He was also Gas Examiner to the Great Western Gas Company, and Chemical Adviser to the Trinity House. We have in these pages frequently pointed to the merits of his various text-books of practical chemistry, and within the last few days he corrected the final proofs of a new work which will shortly be published. His success as a teacher and his skill as an analyst are so well known that they need not be alluded to further, and at present we can only take this, the earliest, opportunity of expressing the regret that will be so widely felt at his loss. Unfortunately, he leaves a widow and a family without provision, and the labours of the committee to which we referred last week will be continued for their benefit. The Hon. Treasurer is Mr. F. W. Bayley, Royal Mint, E., who will gladly acknowledge any contributions that may be sent to him.

New Reagent for Carbohc Acid.—One to two drops of the liquid should be placed in a porcelain capsule with the addition of 2 or 3 drops of a solution of molybdic acid in 10 parts of sulphuric acid. If carbohc acid is present there is produced immediately a light yellow or yellowish brown colour, which changes to a maroon, and finally to a purple. The solution should be very dilute.—*Polyt. Notisblatt.*

CORRESPONDENCE.

PRECIPITATION OF METALLIC COPPER BY POOR COPPER MATT.

To the Editor of the Chemical News.

SIR,—Having lately read the very interesting paper by Mr. Dixon, on the "Method of Extracting Gold, Silver, and other Metals from Pyrites," which appeared in the *CHEMICAL NEWS*, I should like to correct one statement made there in reference to the precipitation of metallic copper, from solution of its sulphate, by poor copper matt. In the above paper Mr. Dixon says (*CHEMICAL NEWS*, vol. xxxviii., p. 303) that:—"Whilst experimenting on the removal of copper from solution I found that this could be conveniently done by filtering the slightly acid solution by ground matt obtained from the same ore by simple melting. This method of separating copper from solution may be of advantage in treating poor copper ores or pyrites containing small quantities of copper:—The matt obtained by simply melting poor cupreous pyrites, with the addition of sufficient roasted ore to form a flux for the silica present, consists of sulphide of iron containing more or less sulphide of copper; and by filtering through a bed of this matt, the solution of sulphate of copper obtained by calcining and extracting a large portion of the ore, the copper is deposited, and the iron goes into solution. I expected to find that the whole of the iron could be thus removed from the matt which would be converted into sulphide of copper, but found that in all cases the action stopped short of this. The percentage of copper in the treated matt varied from 30 to 33 per cent, approximating, therefore, to copper pyrites, which contains 34.6 per cent of copper. From this residue refined copper could be made in three operations."

Now, before having read that paper, I had already been making experiments on the same subject to see if it were possible to raise the percentage of copper in poor matt by some simply wet process, in which the use of acids could be avoided, as by calcining poor cupreous pyrites in heaps, by which means large quantities of sulphates of copper and iron would be produced. I found by filtering a solution of sulphate of copper through a bed of matt containing from 13 to 16 per cent copper, smelted in cupola furnaces, that the percentage of copper was nearly doubled, but on examination I found that this was accounted for by the presence of a large quantity of metallic iron which had been reduced in the furnaces, and had been tapped out with the matt. This iron was disseminated through the mass in very minute globules, and was easily extracted from the ground matt by means of a magnet. To make sure that it was only owing to the presence of this metallic iron that precipitation of copper took place I took some samples of poor matt produced in reverberatory furnaces, and found that no precipitation of copper whatever took place, except in one instance, where 0.83 per cent of copper was thrown down by a sample of matt containing 22 per cent copper, and in this case a small quantity of metallic iron was detected by the magnet, which may possibly have been derived from pounding the sample, which was very hard, in a rather soft iron mortar. The following is the result obtained by treating various samples of poor matt containing metallic iron with a sulphate of copper solution.

A saturated solution of cupric sulphate, slightly acidified (as recommended by Mr. Dixon), was filtered through a layer of 1000 grains finely ground matt, placed in a funnel, the neck of which was plugged with asbestos, till no further precipitation of copper took place. The matt was then washed, dried, and the copper estimated.

	I.	II.
	Per cent.	Per cent.
Matt, after above treatment = Copper ..	26.50	25.63
" before " " = " ..	14.35	14.12
Copper thrown down from solution ..	12.15	11.51

No. I. was a mixture of samples from about 250 taps, and No. II. from above 1200 taps.

The analysis of the two samples of matt is as follows:—

	I.	II.
Copper	14'35	14'12
Iron	56'14	55'47
Sulphur	29'51	30'41
	100'00	100'00
or—		
Sulphides of copper and iron	89'06	88'69
Metallic iron	10'94	11'31
	100'00	100'00

It will be thus seen that the quantity of copper precipitated is directly proportional to the quantity of metallic iron contained in the matt, the theoretical amount of iron required to precipitate copper from solution being in the proportion of 1 part iron to 1'133 parts copper; the amount of copper precipitated in the above cases being in the proportion of:—

No. I.	Iron.	Copper.
No. II.	I to	1'110
	I to	1'017

This proportion is far higher than the results generally obtained by precipitating copper from solution by means of scrap iron, the quantity used in practice ranging from 2 to 4 times the amount required by theory.

The precipitation of copper by the matt was very rapid, owing to the very fine state of division of the metallic iron. This reduction of iron to the metallic state is one great drawback to the smelting of poor cupreous pyrites in cupola furnaces, as unless a very strong and hot blast is kept up (in order to run the ore down as rapidly as possible) the iron soon forms and sets below the tuyeres commencing at the back of the furnace and gradually extending till the tap holes are quite choked up with it; the furnace has then to be cooled down and the mass of iron cut out.

As the above results may be of interest to some of your readers, and may correct some erroneous impressions on this subject, I should feel obliged by your inserting this letter.—I am, &c.,

A. E. BARCLAY.

Bells Cove Copper Mines, Newfoundland,
April 17, 1879.

THE VITRIOL MANUFACTURE,

To the Editor of the Chemical News.

SIR,—My name occurring so many times in Dr. Hurter's letter (CHEM. NEWS, vol. xxxix., p. 170) calls for remark; but first I would mention that Dr. Hurter spells my name incorrectly in every instance.

At the conclusion of his letter he writes—"It is not, in my opinion, of any service to the manufacturer to put him off his guard, and prevent further search by such statements." Now I wish to point out that at the time of reading my three papers before the members of the Faraday Club, I distinctly stated that the instances E, F, and G (working with towers) were not satisfactory, and I urged the members of the Club to work in this direction. With regard to the oxidation of the arsenious acid by the nitrous anhydride, this reaction was repeatedly tried in the Laboratory without success; still, finding nitric oxide in the outlet, and also finding that the arsenious acid was oxidised to arsenic acid in running down the tower, I supposed that as the physical conditions were different the supposed reaction might take place in the tower. When I perceived I might have been mistaken, and that N_2O_4 might have been the oxidiser, and not N_2O_3 , I wrote the letter (CHEM. NEWS, vol. xxxvii., p. 195).

Dr. Hurter, in his letter already quoted, and also in a statement made to the Noxious Vapours Committee of

the Alkali Makers' Association, asserts that the exit gases contain sufficient oxygen to convert the N_2O_4 into N_2O_3 , and therefore that N_2O_4 is never to be found in exit gases.

It would be very interesting if those chemists who have observed N_2O_4 passing in company with oxygen would come forward and state what they have found.

I have a set of experiments going on which may tend to throw some light on the subject of the oxidation of the arsenic. I still maintain that N_2O_4 does go away in the exit gases when from 4 to 8 per cent of oxygen is present. This may be only under certain conditions of working, but still I have found it, and so have others.

I never considered that the samples from absorber exits E, F, and G (CHEM. NEWS, vol. xxxvii., p. 157) were satisfactory solutions of the question. E gives more nitre escaping than used; of F I gave no exit tests; and G I stated that the chambers were old, much air was drawn in at the last ones, and the amount of nitre unaccounted for was 20 per cent.

Dr. Hurter states that my figures prove that the chemical losses in the chambers are from 10 to 15 per cent. Now I stated distinctly that in samples A, B, C, and D the total unaccounted for was 5 to 9 per cent. This quantity might be decomposed in the chambers where steam is present, but I have no proof of it, neither do I know any one who has.

Dr. Hurter draws his conclusions on the exit escape of nitre from the averages of seven different works. I am convinced that as long as this is done we shall never arrive at the true cause of all the nitre losses. These losses must be studied in each individual works, and where the conditions always remain the same. One works may have an equal number of chambers of a similar size with another works, doing the same work, and with denitrators and absorbers equal; one may be using 1'8 per cent of nitre, and the other 3 per cent; and again, in the same works an even number of chambers may be divided into two sets, and yet one may always consume more nitre than the other, in order to prevent an escape of SO_2 .

The difficulties which surround this subject may be inferred from Mr. E. Jackson's letter (CHEM. NEWS, vol. xxxviii., p. 147), in which he states—"Thus during the past week the average quantity of nitre found in the exit gases does not exceed 4 lbs. per ton of stone, while about a month ago the quantity found during one week exceeded the quantity used for that week." Since then Mr. Jackson has informed me that during one whole period of six months he could account for all the nitre used, within a small percentage, without reference to any theory of reduction to nitrogen or nitrous oxide. Perhaps the following experiments may be of use to investigators:—

1. The acid was issuing from a Glover tower at 150° Tw. cold, and at a temperature of 290° F. containing 0'02 per cent of nitrogen compounds, calculated at N_2O_3 by the mercury method. A damper was now inserted in the iron pipe (leading the SO_2 gas into the tower) having in its centre a 4-inch circular opening, a direct (extra) way being opened for the gases to go direct into the chambers. The small quantity of SO_2 which now went up the tower effected no concentration, the acid was perfectly denitrated, the issuing acid was from 90° to 100° F., and no saving of nitre was effected.

2. The nitrous vapours were being boiled off in pots placed in a very cool part of the flue leading from the kilns to the chambers (working without towers). The consumption of nitre was 4'2 per cent upon 47 per cent of sulphur. The potting place was changed to a bright red-hot portion of the flue; this change caused no more nitre to be used, nor was the colour of the chambers or escape in any way affected. The potting place was again changed to where the active combustion was going on, and this spot was nearly at a white-heat: within a few hours from this change the chambers began to grow pale, hard potting was necessary, more nitre was used for some time—to try

and bring the colour of their chambers to their normal conditions—without success, and when the first and original potting place was resumed the nitre went down to its original amount.

3. It is thought by some that the last chambers reduce the N_2O_4 to NO_2 by means of the excess of water present (i. e., the low density of the acid). It was tried to work the last chambers stronger than usual, viz., 136°Tw. , to 140°Tw. ; the result was a greater use of nitre than before, even taking into consideration the amount absorbed by the acid. This experiment was going on for many months, and during the whole of the time the total acidity of the exit was more regular and lower than when the last chambers were worked weaker.

4. Tests of inlet and outlet of absorbing towers:—

a = Na_2CO_3 neutralised by S compounds, grains per cubic feet.

b = Na_2CO_3 neutralised by N compounds, grains per cubic feet.

c = oxygen per cent by volume.

		Inlet.	Outlet.
1.	a.	5'90	3'65
	b.	1'40	0'70
	c.	9'80	Colourless.
2.	a.	2'98	2'79
	b.	1'02	0'08
	c.	10'25	Colour pale straw.
3.	a.	2'55	2'36
	b.	1'79	0'11
	c.	11'25	Colour straw.
4.	a.	4'80	4'05
	b.	1'69	1'11
	c.	5'90	Colourless.
5.	a.	3'62	3'42
	b.	1'84	1'02
	c.	6'60	Colourless.
6.	a.	1'07	1'00
	b.	2'04	0'11
	c.	10'27	Colour straw.

Before leaving this subject I wish to point out that Dr. Hurter has applied the most refined methods to the ascertaining of the mechanical losses; he should therefore be able to give us them within 1 or 2 per cent.

I have heard it remarked that Lunge has stated that my process for the estimation of nitrous compounds in nitrous vitriol, by shaking up with mercury and measurement of the nitric oxide, is liable to an error of as much as 10 per cent, on account of my not taking the temperature and pressure into consideration. I have heard that this has been abstracted into the *Journal of the Chemical Society*, though I have not seen it,—therefore it is necessary for me to quote from my paper (CHEM. NEWS, vol. xxxvii., p. 45):—"For technical purposes this volume will be found accurate enough, but in cases where extreme accuracy is required the tube must be left to itself for several hours, the temperature and pressure noted, and the necessary corrections made."—I am, &c.,

GEORGE E. DAVIS.

Heaton Chapel, Stockport,
April 28, 1879.

Determination of Nitric Acid Contained in Commercial Lime-Juice.—F. Dotto-Scribani.—The author takes 100 c.c., boils the liquid, and neutralises exactly with baryta water. Citrate of barium is formed and precipitated, whilst barium nitrate remains in solution. The liquid is let cool, the citrate is separated from the nitrate by filtration, and the latter salt is decomposed with sulphuric acid. The barium sulphate obtained is filtered, washed, &c., and weighed with the usual precautions, and on multiplying its weight, less the ash of the filter, by 0'4635 the quantity of nitric acid present in 100 c.c. of the juice is obtained.—*Gazzetta Chimica Italiana*.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 62, February, 1879.

Report Presented by M. le Comte du Moncel on Behalf of the Committee of Economical Arts on M. E. Reynier's Electric Light Regulator.—The committee decide that M. Reynier has solved the problem in question. His arrangement has been successfully tried at the station of the Northern railway and in M. Breguet's work-shop, where it has acted for some hours with regularity under the influence of the current of a small Gramme machine. The arrangement is figured on the accompanying plate.

Report Presented by M. Aimé Girard on M. Kuhlmann's (junior) Methods for Recognising the Composition and Measuring the Volume of the Gases and Acid Vapours given off by the Chimneys of Chemical Works.—By means of this arrangement a manufacturer can at any hour inform himself concerning the nature of the gases traversing his chimney. To establish the composition of the chimney-gases and to detect, e.g., the presence of hydrochloric acid therein, M. Kuhlmann introduces into the shaft a glass tube, which communicates with a series of test-tubes, at the end of which is a cistern of water, which acts as a respirator. The flow of water compels the gas to traverse the series of test-tubes, the first of which contains caustic soda coloured with litmus. Then follow solutions of barium chloride and silver nitrate. The dimensions of the cistern are such that five or six hours are required for the total efflux of the water which it contains, and the strength of the alkaline liquid coloured blue is such that in the regular course of work five or six hours are required to turn them red. An earlier colouration gives immediate warning. M. Kuhlmann's method of measuring the total volume of gases escaping is as follows:—When he wishes to know the speed of the gases he introduces at the bottom of the chimney a certain volume of a strong-coloured gas, such as hyponitric acid, which accompanies the colourless gases up the chimney and presents itself along with them at the top. Knowing, then, the volume of the chimney and the mean temperature it is possible to calculate from the time which has elapsed the quantity of gas emitted.

Bulletin de la Société Chimique de Paris,

No. 6, March 20, 1879.

Action of Mono-chlor-acetic Acid upon Sulphocyanic Acid and its Salts.—M. Nencki.—On heating a mixture of 1 mol. chlor-acetic acid dissolved in double its weight of water with 3 mols. ammoniac sulphocyanate, the liquid obtained deposits crystals of a new compound, rhodanic acid, $\text{C}_3\text{H}_2\text{NS}_2\text{O}$, which crystallises in tables and six-sided prisms of a fine yellow, scarcely soluble in cold water, but readily in alcohol, ether, and alkalis. It melts about 168° to 170° with partial decomposition, yielding a brown-red liquid. Under the influence of feeble oxidising agents it produces colouring matters. A red and a violet matter have thus been obtained which in acid solutions dye silk and wool, and in alkaline solutions produce a blue shade upon cotton.—*Journal f. Praktische Chemie*, xvi., 1.

Action of Cyanogen upon Albumen.—O. Loew.—By treating albumen with cyanogen gas the author obtains three new compounds, cyalbidin, oxamoidin, and a third body not named.—*Journ. f. Prakt. Chemie*, xvi., 60.

Gazzetta Chimica Italiana.
Anno viii., 1878. Fasc. x.

Certain Derivatives of Campho-thymol.—E. Paterno and F. Canzoneri.—An account of nitroso-campho-thymol and amidico-campho-thymol.

Constitution of the Cymen of Cuminic Alcohol and of its Thymols.—E. Paterno and P. Spica.—Not suitable for abstraction.

Propyl-benzoic Acid.—E. Paterno.—A claim of priority as against H. Koerner.

Benzylated Cymen.—Dr. Girolamo Mazzara.—The author finds this compound to consist of—Carbon, 91.00; hydrogen, 9.10; and gives it the formula—



Process for Discovering the Effective Quantity of Tartaric Acid Contained in Dregs of Wine.—Prof. Francesco Dotto-Scribani.—This process serves for determining the effective quantity of tartaric acid not merely in unadulterated but even in the dregs of plastered wines. 10 grms. of the dregs are reduced to a very fine powder and dried in the water-bath, and then moistened and made into a paste with pure hydrochloric acid in a porcelain capsule, and allowed to stand for twenty-four hours. The mass is then lixiviated with boiling distilled water, the liquid is filtered, and this treatment is repeated with the residue upon the filter till the liquid passing through no longer reddens litmus paper. The filtrate is then boiled in a porcelain capsule, and whilst it is continually stirred with a glass spoon milk of lime obtained by levigation and passed through a strainer is added little by little until the liquid begins to turn red litmus paper blue. The capsule is then allowed to cool and the contents thrown upon a double filter, which has been previously well washed, and the precipitate is then washed, until on acidulating the washings with nitric acid and testing with nitrate of silver no turbidity is observed. The precipitate with the double filter is then dried in the water-bath and weighed, having care first to detach the outer from the inner filter, and to place the former in the weight-scale, and from its weight is deduced the effective quantity of tartaric acid, since 100 parts of calcium tartrate contain 57.69 parts of tartaric acid. This process is founded on the following principles:—The acid tartrate of potassium and the tartrate of calcium are less soluble in boiling water than in such water acidulated with hydrochloric acid, with which the two salts produce chlorides of potassium and calcium, while tartaric acid is set free. The colouring matter of the dregs is insoluble in hydrochloric acid. Milk of lime has no action upon the chlorides, but transforms the tartaric acid into calcium tartrate.

MISCELLANEOUS.

Chemistry in Australia.—Mr. R. W. E. MacIvor A.I.C., &c., late Senior Assistant to Professor Dittmar, at Anderson's College, Glasgow, has for the last three years been engaged in lecturing to the farmers throughout Victoria on Agricultural Chemistry, and has succeeded in impressing upon them the importance of the subject, for many come 40 and even 50 miles to listen to him. In referring to Mr. MacIvor's discourses, the *Lancefield Chronicle* for March 20 says:—"The amount of good derived from these scientific dissertations has been incalculable to the agriculturist, and the knowledge imparted has increased the prosperity of the country." It is but fair to mention that these lectures are given free of charge to the agricultural community, the expense attending their delivery being borne by the Hon. William J. Clarke, M.L.C., President of the forthcoming Melbourne International Exhibition.

Detection of Arsenic.—The matter containing the arsenious or arsenic acid is introduced into a Marsh's apparatus, and mixed with a concentrated solution of caustic potassa and a little aluminium foil. Arseniuretted hydrogen is disengaged on applying heat. Antimoniu-retted hydrogen is not formed.—*Moniteur Scientifique*.

Technological Examinations.—The Council of the Society of Arts, having received an application from the City and Guilds of London Institute for the Advancement of Technical Education, offering to take charge of the Technological Examinations established by this Society in 1873, and carried on up to the present time, have resolved to transfer these examinations to the charge of the Institute. The Council have also ascertained that the Science and Art Department will assist the City Institute in conducting the examinations, in the same way as it has hitherto assisted the Society of Arts. The Technological Examinations for the present year will, therefore, be carried on under the direction of the Institute, and all communications on the subject should be addressed to the Hon. Secretaries, City and Guilds of London Institute, Mercers' Hall, E.C.

Science-Teaching in Public Schools.—A correspondent suggests that a careful and trustworthy account of the teaching of science in schools would be of great value, by laying before the public facts and not mere theories. As an instance of such theories he mentions the assumption of no less eminent a man than Sir J. Lubbock that boys would be anxious or willing to study science in play hours. He points out from among 500 boys few care to hear even such a lecturer as Prof. Tyndall, and none would go a second time except other work was remitted in compensation. Our friend informs us that at Harrow and at some other public schools any boy at the desire of his parents can give several hours weekly to practical work in chemistry or physics. "The parents of boys in general do not care about their learning practically, and it is from them, rather than from the masters, that the difficulty arises." May we suggest that such indifference or disinclination on the part of parents springs from the fact that successful practical work in science counts for little in determining a boy's standing in the schools?

MEETINGS FOR THE WEEK.

- MONDAY, 12th.—Royal Geographical, 8.30.
Society of Arts, 8. "Recent Advances in Telegraphy," by W. H. Preece. (Cantor Lectures.)
- TUESDAY, 13th.—Civil Engineers, 8.
Royal Institution, 3. "The Intellectual Movement of Germany from the Middle of the Last to the Middle of the Present Century," Prof. Hillebrand.
Photographic, 8.
Anthropological, 8.
- WEDNESDAY, 14th.—Society of Arts, 8. "The Automatic Hydraulic Brake," Mr. E. D. Barker.
Geological, 8.
Microscopical, 8.
- THURSDAY, 15th.—Royal, 8.30.
Royal Institution, 3. "Dissociation," by Prof. Dewar.
Royal Society Club, 6.30.
Society of Arts, 8. "The History of Alizarin and Allied Colouring Matters, and their Production from Coal-Tar," by W. H. Perkin, F.R.S.
Chemical, 8. "On Nitric Acid, Part II.," Robert Warington. "On Alkaloids of the Veratrum, Part III.," Dr. Wright and Mr. Luff. "On Alkaloids of the Veratrum, Part IV.," Dr. Wright. "On Alkaloids of the Aconite, Part IV.," and on Japanese Aconite Roots, Dr. Wright and Mr. Luff. "Action of Hydrochloric Acid on Manganese Dioxide," S. Pickering. "Composition of Milk in Health and Disease," A. W. Elyth. "Notes on the Effect of Alcohol on the Chemistry of Indigestion," W. H. Watson.
- FRIDAY, 16th.—Royal Institution, 9. "Etude Optique de l'Elasticité," Prof. Cornu.
- SATURDAY, 17th.—Royal Institution, 3. "Architecture," by Mr. H. H. Statham.

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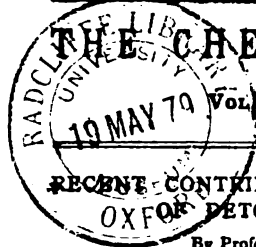
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Vol. XXXIX. No. 1016.

RECENT CONTRIBUTIONS TO THE HISTORY
OF DETONATING AGENTS.*

By Professor ABEL, C.B. F.R.S.

(Concluded from p. 200.)

In comparing the effects of these nitro-glycerin preparations with each other and with compressed gun-cotton and preparations of it, by detonating equal quantities quite unconfined upon iron plates, the results appear to establish great superiority, in point of violence of action, or destructive effect, of the more rigid explosive agents (the gun-cotton preparations). Thus, employing iron plates 1 inch thick (supported upon an anvil with a central cavity), and 4 ozs. of each material unconfined, the charges being all about the same diameter, exploded by detonators of equal strength, and simply resting upon the upper surface of the plate, compressed gun-cotton produced a considerable indentation of the upper surface of the plate, and long cracks in the lower surface; a species of nitrated gun-cotton, called tonite, produced a much shallower indentation, though still a very marked one, but did not crack the lower surface. Dynamite produced only a very slight impression upon the plate, and none could be detected by the eye on the plate upon which the blasting gelatin was exploded. The difficulties, brought out by past experience, which attend the contrivance of really comparative tests of the explosive power of such substances as those under discussion, is well exemplified by the foregoing results, which were influenced to the maximum extent by the physical characters of the several substances when applied, as they were upon these iron plates, in a perfectly unconfined condition, so that the particles were free to yield to the force of the initiative detonation in proportion to their mobility. But, for this very reason, these experiments afford excellent illustration of the extent to which the development of detonation and the sharpness of its transmission through the mass is influenced not only by the inherent sensitiveness of the substance to detonation (or by its chemical instability) but also by the degree of proneness of their particles to yield mechanically to the force of a blow as applied by an initiative detonation. Thus, although in comparing two substances of similar physical characters, compressed gun-cotton and compressed nitrated gun-cotton or tonite, the superiority of the pure compound over the mixture, in point of sharpness and violence of action, is well illustrated, a comparison of the result furnished by the weakest of the four explosive agents tried, viz. tonite, with that of the substance which should be superior to all the others in explosive force (*i.e.* the blasting gelatin) demonstrates the important influence which the comparatively great rigidity of the mass in the one case exerts in favouring the completeness and sharpness of its detonation in open air, and the great disadvantage under which the other explosive is applied, arising out of the plastic and therefore readily yielding nature of the material. But if, by exposure to a moderate degree of cold, this plastic nitro-glycerin preparation is made to freeze (for it partakes of the property of the liquid itself of freezing at a temperature above the freezing-point of water, and becomes thereby converted into at least as rigid a substance as the two descriptions of gun-cotton) its detonation upon an iron plate produces an indentation, as well as a destructive effect upon the lower surface of the plate, very decidedly greater than those furnished by

the corresponding amount of pure compressed gun-cotton. Similarly, the effect produced by the detonation of dynamite upon a plate of the kind used, is but little inferior to that of gun-cotton, and decidedly greater than that of tonite, if it is employed in the frozen condition.

A series of experiments has been made with cylinders of lead having a central perforation 1.3 inches in diameter extending to a depth of 7 inches and leaving solid metal beneath of a thickness ranging from 3.5 to 5.5 inches, according to the size of the cylinders used. These furnished results of considerable interest as illustrating the action of these several detonating agents. Charges of 1.25 ozs. of each explosive substance were used throughout the experiments, and were placed at the bottom of the holes. By the detonation of the charges the cylindrical holes in the lead were enlarged into cavities of a pear shape (and sometimes approaching the spherical form), of various diameters; in some instances the metal was besides partially torn open in a line from the bottom of the charge-hole to the circumference of the lower face of the cylinder; and in the case of some of the gun-cotton charges, the fissure in the metal in this direction was complete, the base of the block being separated from the remainder, in the form of a cone. In the first place the portions of the holes above the charges were simply left open; in the subsequent experiments they were filled up to a level with the upper surface, with dry, fine, loose sand, or with water. The dimensions of the cylinders were increased in successive experiments until, in the case of every one of the explosives used, the mass of metal was sufficiently great to resist actual fracture at the base of the cylinder. Under the conditions of these experiments, more or less considerable resistance being opposed to the mechanical dispersion of the plastic explosive substances, their detonation was greatly facilitated, though even then the holes in the lead blocks being left open to the air, some amount of the blasting gelatin evidently escaped detonation; the widening of the upper part of the charge-hole in experiments of this nature made with the gelatin indicated that detonation was transmitted to small portions dispersed in the first instance and in the act of escaping from the block. In all the experiments, whether the holes were left open or filled with sand or water, the effect produced upon the base of the block by the detonation of compressed gun-cotton was considerably more violent than with the other explosive agents, indicating a sharpness of action which was only shared by the blasting gelatin when used in a frozen state in one of these experiments. The dimensions of the cavities produced by the gelatin were, at the largest part, considerably greater than those produced by the dynamite and nitrated gun-cotton (tonite), and slightly greater than those of the gun-cotton charges; but in the latter, the fracture of the base of the cylinder gave rise in most of the experiments to an escape of force, so that in these cases the effects of the detonation could not be well compared by measurements of the cavities. When the gelatin was converted by freezing into a rigid mass its superiority in explosive force even over compressed gun-cotton was well illustrated; the base of the lead block was all but blown out, the cavity produced was considerably the largest, and the suddenness and violence with which motion was imparted to the water tamping caused the top of the block also to be blown off in the form of a cone. An excellent illustration was obtained by comparing this result with those furnished by the gelatin in its normal plastic state of the influence exercised by the physical condition of an explosive substance upon the rapidity and completeness with which detonation is transmitted through its mass.

The difficulties attending the application of blasting gelatin, in some directions in which explosive agents are applied, on account of the uncertainty attending the development of its explosive force, even with the use of a comparatively powerful detonator, unless it be very strongly confined, has led to attempts to reduce its non-sensitive-

* Abstract of a Paper read before the Royal Institution of Great Britain, Friday, March 21, 1899.

ness to detonation by mixing it with materials intended to operate either by virtue of their comparatively great sensitiveness, or of their property, as solids, of reducing the very yielding character of the substance, or in both ways.

Some of these attempts have been attended with considerable success. Thus the incorporation of about 10 per cent of the most explosive form of gun-cotton or trinitro-cellulose, in a very finely divided state, with the gelatin, renders it so much more sensitive that it can be detonated with certainty in the open air by means of the strongest detonating cap now used for exploding dynamite. This effect appears to be less due to the comparative sensitiveness of gun-cotton to detonation than to the modification effected in the consistency of the material, which, though still plastic, offers decidedly greater resistance to a blow than the original gummy substance. The particles of hollow fibre of the gun-cotton appear also to have the effect of absorbing small quantities of nitro-glycerin which are less perfectly united with the soluble gun-cotton than the remainder, and which, existing as they do in somewhat variable proportions in the gelatin, have occasionally an objectionable tendency to exudation, if the incorporation of the ingredients has been less perfect than usual. The substance, when modified as described, has no longer that great adhesiveness which is exhibited by it in the original state, and which renders it less easy to manipulate.

Lastly, its explosive force appears to be in no way diminished by this modification of its composition, on the contrary, its superiority in this respect to compressed gun-cotton becomes more manifest, as demonstrated by some of the experiments with lead blocks, while its action partakes of that sharpness peculiar to the detonation of the rigid gun-cotton, as indicated by the fissure of that part of the metal situated beneath the charge. Finely divided cotton fibre has a similar effect to trinitro-cellulose in modifying the physical character and increasing the sensitiveness to detonation of the blasting gelatin, but its explosive force is, of course, proportionately reduced with its dilution with an inert substance.

Nobel has made the interesting observation that an addition to the blasting gelatin of small proportions of certain substances rich in carbon and hydrogen, which are soluble in nitro-glycerin, such as benzol and nitro-benzol, increases to a remarkable extent the non-sensitiveness to detonation of the original material; and this observation has led to experiments being conducted by engineer officers in Austria, with a view of endeavouring to convert the blasting gelatin into a material which should compete, as regards some special advantages in point of safety, with wet gun-cotton in its application to military and naval purposes, and especially as regards non-liability to detonation or explosion by the impact of rifle bullets. If boxes containing dry compressed gun-cotton are fired into from small arms even at a short range, the gun-cotton is generally inflamed, but has never been known to explode; the sharpness of the blow essential to the latter result, which the bullet might otherwise give, being diminished by its penetration through the side of the box before reaching the explosive. It is scarcely necessary to state that wet gun-cotton, containing even as little as 15 per cent of water, is never inflamed under these conditions. On the other hand, dynamite is invariably detonated when struck by a bullet on passing through the side of the box, and blasting gelatin, though so much less sensitive than dynamite, behaves in the same way in its ordinary as well as in the frozen condition. The Austrian experiments indicated that the gelatin when intimately mixed with only 1 per cent of camphor, generally, though not invariably, escaped explosion by the impact of a bullet, but that when the proportion of camphor amounted to 4 per cent the material was neither exploded nor inflamed, though, in the frozen state, it was still liable to occasional explosion. These results were considered indicative of a degree of safety in regard

to service exigences, approaching that of wet compressed gun-cotton. The camphoretted gelatin still labours, however, under the disadvantage of being readily inflammable, and of burning fiercely, and consequently of giving rise, like dynamite and dry gun-cotton, to violent explosion or detonation, if burned in considerable bulk; a result which was explained by the lecturer in his discourse delivered at the Royal Institution in 1872. Moreover, the camphoretted blasting gelatin is so difficult of detonation by the means ordinarily applied, that a large initiative charge of a very violent detonating mixture consisting of pure specially prepared trinitro-cellulose and nitro-glycerin is prescribed by the Austrian experimenters as being indispensable to its proper detonation.

The action of camphor and of other substances rich in carbon and hydrogen in reducing greatly the sensitiveness to detonation of the preparation of soluble gun-cotton and nitro-glycerin, is not to be traced to any physical modification of that material produced by the addition of such substances, and no satisfactory theory can at present be advanced to account for it on chemical grounds.

The camphoretted gelatin, like Nobel's original gelatin itself, may be kept immersed in water for a considerable time without any important change; the surface of the mass thus immersed becomes white and opaque, apparently in consequence of some small absorption of water, but no nitro-glycerin is separated, and on re-exposure to the air the material gradually assumes once more its original aspect. It has consequently been proposed to render the storage of blasting gelatin comparatively safe by keeping it immersed in water till required for use, but the test of time is still needed to establish the unalterableness of the material under this condition.

There can be little question that this interesting nitro-glycerin preparation, either in its most simple form or modified in various ways by the addition of other ingredients, promises, by virtue of its great peculiarities as a detonating agent, to present means for importantly extending the application of nitro-glycerin to industrial purposes; and it is not improbable that through its agency this most violent of all explosive agents at present producible upon a large scale may also come to acquire special value for important war purposes.

It has been pointed out that the complete solidification by freezing of plastic preparations containing nitro-glycerin, such as dynamite and the blasting gelatin, has the effect of facilitating the transmission of detonation throughout the mass, and of thus developing or increasing the violence of their action, under certain conditions of their applications, *i.e.*, when they are either freely exposed to air or not very closely or rigidly confined. But while, under circumstances favourable to the detonation of these substances, when in the frozen state, their full explosive force is thus much more readily applied than when they are in their normal (thawed) condition, the frozen substances are less sensitive to detonation by a blow or an initiative detonation. On the other hand, if subjected to the rapid application of great heat (as, for example, by the burning of portions of a mass of the explosive substance itself), a detonation is much more readily brought about with the frozen material than if it be in its normal condition. Thus a package containing 50 lbs. of cartridges of plastic dynamite when surrounded by fire burned away without any indication of explosive action; on submitting 10 lbs. of frozen dynamite to the same treatment that quantity also burned without explosion, though at one time the combustion was so fierce as to indicate an approach to explosive action; but when the experiment was repeated on the same occasion with 15 lbs. frozen dynamite a very violent detonation took place after the material had been burning for a short time, a deep crater being produced in the ground beneath.

The following is offered as the most probable explanation of this result. When a mass of dynamite, as in these cartridges, is exposed to sufficient cold to cause the nitro-glycerin to freeze, it does not become uniformly

hardened throughout, partly because of slight variations in the proportion of nitro-glycerin in different portions of the mixture composing the cartridge, and partly because unless the exposure to cold be very prolonged the external portions of the cartridges will be frozen harder or more thoroughly than the interior. It may thus arise that portions of only partially frozen or still unfrozen dynamite may be more or less completely enclosed in hard crusts, or strong envelopes, of perfectly frozen and comparatively very cold dynamite. On exposure of such cartridges to a fierce heat very rapidly applied, as would result from the burning of a considerable quantity of the material, some portion of one or other of the cartridges would be likely to be much more readily raised to the igniting or exploding point than the remaining more perfectly frozen part in which it is partly or completely imbedded. If the ignition of this portion be brought about (which it will be with a rapidity proportionate to the intensity of heat to which the cartridge is exposed), the envelope of hard frozen dynamite by which it is still more or less completely surrounded and strongly confined, will operate like the metal envelope of a detonator, in developing the initial pressure essential for the sudden raising of the more readily inflammable portion of the dynamite to the temperature necessary for the sudden transformation of the nitro-glycerin into gas, and will thus bring about the detonation of a portion of the cartridge, which will act as the initiative detonator to the remainder of the dynamite. On igniting separately, at one of their extremities, some dynamite cartridges which had been buried in snow for a considerable period, the lecturer has observed that, as the frozen material gradually burned away, very slight but sharp explosions (like the snapping of a small percussion cap on a gun nipple) occurred from time to time, portions of the frozen dynamite being scattered with some violence. He did not succeed in obtaining actual detonation by thus burning frozen cartridges surrounded by others in a similar condition, but he has been informed by Mr. McRoberts, of the Ardeer Dynamite Works, that he has more than once detonated a small heap of hard-frozen cartridges, weighing altogether 1 lb., by igniting one cartridge which was surrounded by the remainder. These facts appear to substantiate the correctness of the foregoing explanation. They point to the danger of assuming that, because dynamite in the frozen state is less sensitive to the effects of a blow or initiative detonation than the thawed material, it may therefore be submitted without special care to the action of heat, for the purpose of thawing it. Instances of the detonation, with disastrous results, of even single cartridges of frozen dynamite, through the incautious application of considerable heat (as, for example, by placing them in an oven, or close to a fire), have been, and are still, of not unfrequent occurrence, even though Mr. Nobel has insisted upon the application of heat through the agency only of warm water, as the sole reliable method of safely thawing dynamite cartridges.

While the sensitiveness to detonation of air-dry gun-cotton remains unaffected by great reduction in temperature of the mass, and while in this respect it presents advantages over nitro-glycerin preparations, wet gun-cotton becomes very decidedly more susceptible to detonation when frozen. Thus the detonation of gun-cotton containing an addition of from 10 to 12 per cent of water is somewhat uncertain with the employment of 100 grs. of strongly-confined fulminate, and 200 grs. are required for the detonation of the substance when containing 15 to 17 per cent of water; but the latter in a frozen state can be detonated by means of 30 grs. of fulminate, and 15 grs. are just upon the margin of the amount requisite for detonating with certainty frozen gun-cotton containing 10 to 12 per cent of water. The deadening effect of solid water upon the sensitiveness of gun-cotton to detonation is, in fact, intermediate between those of a liquid and of inert solid substances.

The effects produced and products formed by the explo-

sion of gun-cotton in perfectly closed spaces, both in the loose and the compressed form, and by its detonation in the dry and the wet state, have been made the subject of study by Captain Noble and Mr. Abel, the method of research pursued being the same as that followed in their published researches on fired gunpowder; results of considerable interest in regard to the heat of explosion, the pressures developed, and the products of explosion of dry and wet gun-cotton have been obtained, which are about to be communicated to the Royal Society.

It may briefly be stated that the temperature of explosion of gun-cotton is more than double that of gunpowder (being about 4400° C.); that the tension of the products of explosion, assuming the material to fill entirely the space in which it is fired, is considerably more than double that of the powder-products under the same conditions; that the products obtained by the explosion of dry gun-cotton are comparatively simple and very uniform under different conditions as regards pressure; that the products of *detonation* of dry gun-cotton do not differ materially from those of its explosion in a confined space, but that those furnished by the detonation of *wet* gun-cotton present some interesting points of difference. Messrs. Noble and Abel are extending their investigations to the nitro-glycerin preparations.

The great advance which has been made within the last twelve years in our knowledge of the conditions which determine the character of the metamorphosis that explosive substances undergo, and which develop or control the violence of their action, finds its parallel in the progress which has been made in the production, perfection, and application of the two most prominent of modern explosive agents, nitro-glycerin and gun-cotton. Discovered at nearly the same time, less than forty years ago, the one speedily attained great prominence on account of the apparent ease with which it could be prepared and put to practical use; a prominence short-lived, however, because the first, and somewhat rash, attempts to utilise it preceded the acquisition of sound and sufficient knowledge of its nature and properties. Even many years afterwards, when the difficulties attending its employment appeared to have been surmounted, the confidence of its most indefatigable partisans and staunchest friends received a rude shock, from which it needed the support of much faith and some fortitude to recover.

Meanwhile, the other substance, which now shares with it the honours of important victories won over gunpowder continued to be generally regarded as a dangerous chemical curiosity, even for some time after its present position as one of the most important industrial products and useful explosive agents was being gradually but firmly secured for it, step by step, by the talent and untiring energy of a single individual.

Almost from the day of its discovery, the fortunes of gun-cotton continued to fluctuate, and much adversity marked its career, until at last its properties became well understood, and its position as a most formidable explosive agent, applicable on a large scale, with ease, great simplicity, and with a degree of safety far greater than that as yet possessed by any other substance of this class, has now become thoroughly established. Since the lecturer last discoursed on the properties of gun-cotton, seven years ago, this material has attained a firm footing as one of the most formidable agents of defence and offence. For all military engineering operations, and for employment in submarine mines and torpedoes, compressed gun-cotton, stored and used in the wet condition has become the accepted explosive agent in Great Britain; within the last five years upwards of 550 tons have been manufactured for this purpose, and are distributed over our chief naval stations at home and abroad. Germany some years since copied our system of manufacture and use of gun-cotton; France has provided itself with a large supply for the same purposes; and Austria, where the acquisition of bitter experience of the uncer-

tainty of gun-cotton in the earlier stages of history, naturally gave rise to a persistent scepticism regarding its presenting trustworthiness, appears now also about to adopt wet gun-cotton for military and naval uses.

But while the usefulness and great value of compressed gun-cotton in these important directions has been established, its technical application has made but slow progress as compared with that of the simple nitro-glycerin preparation known as dynamite, which, in point of cost of production and convenience for general blasting purposes, can claim superiority over compressed gun-cotton. Already, in 1867, a number of dynamite factories, working under Nobel's supervision, existed in different countries; in that year the total quantity manufactured amounted to 11 tons; in another year the produce had risen to 78 tons; in 1872 it had attained to 1350 tons. Two years afterwards the total production of dynamite was nearly trebled, and in 1878 it amounted to 6140 tons.

There are as many as fifteen factories in different parts of the world (including a very extensive one in Scotland) working under the supervision of Mr. Nobel, the originator of the nitro-glycerin industry, and some six or seven other establishments exist where dynamite or preparations of very similar character are also manufactured.

How far the rate of production of dynamite will be affected by the further development of the value of Nobel's new preparation, the blasting gelatin, it is difficult to foresee, but there appears great prospect of an important future for this very peculiar and interesting detonating agent.

It is hoped that the subjects dealt with in this discourse afford interesting illustration of the intimate connection of scientific research with important practical achievements.—F. A. A.

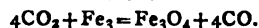
A NEW METHOD OF PRODUCING A COATING OF MAGNETIC OXIDE ON IRON SURFACES.

By GEORGE R. TWEEDIE.

It will be remembered that last May I had the pleasure of laying before the readers of the *CHEMICAL NEWS* a few details respecting Mr. Geo. Bower's process for producing a coating of magnetic oxide of iron by the action of hot air.

A continuation of the experiments has led to the patenting by Mr. Bower of a second process of considerable interest.

It was ascertained, during the time the experiments with the air process were being carried out, that the magnetic oxide was formed by the oxygen of the air combining with the carbon of the iron and forming carbon dioxide, which in presence of the heated iron was split up according to the well-known reaction—



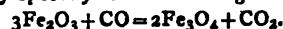
This fact readily explained why the process was unsuitable for wrought-iron or steel.

It was then resolved to carry out a further series of experiments bearing on this fact, and the best means of applying it practically. Pure carbon dioxide prepared in the usual way was first tried. A silvery coating of magnetic oxide was formed after an exposure of about seven hours at a dull red-heat; the coating, however, was crystalline and somewhat brittle, although very hard.

Very long and tedious sets of experiments were then carried out with mixtures of carbon dioxide and air, air and carbonic oxide, and varying exposures of the three together.

The following is the method of procedure now adopted, and which answers most satisfactorily:—The articles are first of all heated, and acted on for a certain period by the products of combustion largely mixed with air from a peculiarly constructed furnace (designed by Mr. Anthony

Bower), burning slack or small coal. In this way a coating of magnetic oxide is formed close to the surface of the iron, but this is often slightly covered with red oxide, Fe_2O_3 . The admission of air to the furnace is then so arranged by a suitable apparatus that a stream of carbonic oxide is passed over the articles for a short time, and the red oxide very speedily reduced to magnetic oxide,—



It is worthy of note that although the red oxide, before reduction to magnetic oxide, is very readily detached from the iron, after the reduction the Fe_3O_4 so formed is perfectly hard and homogeneous. From all appearances it would seem as though a kind of fusion took place, but at the same time it must be recollected that the temperature used (a red-heat) is very low to favour such an adhesion.

The oxide formed by this process has been tested very thoroughly, and withstands the ordinary oxidising influences perfectly.

Mr. Bower has several contracts in hand which are to be dealt with by his processes, and will present opportunities for the practical testing of the two processes from a commercial point of view.

ON THE INFLUENCE OF VARIATIONS OF TEMPERATURE ON THE DEVIATION OF POLARISED LIGHT BY SOLUTIONS OF INVERTED SUGAR.*

By P. CASAMAJOR.

THE researches, of which I propose to give an account in this paper, were suggested by an interesting communication of Dr. Ricketts to this Society, which was presented at our last meeting.

Dr. Ricketts found that the temperature at which the deviation of a solution of inverted sugar becomes 0 is not $90^\circ \text{C}.$, as given by some authors, but 92° , or rather $91.7^\circ \text{C}.$ From this Dr. Ricketts concluded that if a solution of commercial sugar is inverted in the ordinary way, if originally the two constituents of the sugar were cane sugar and inverted sugar, after inversion there must only remain inverted sugar, and if we bring this solution in the saccharometer tube to have the temperature of $92^\circ \text{C}.$, the indication of the saccharometer scale must be 0.

As the inversion of sugar solutions in testing sugars is now almost entirely neglected, it struck me that the introduction of this subject before this Society was of great importance, as it is likely to excite inquiry in this direction, and must lead to interesting discussions which will throw much light on a ground which has not recently been explored. I hope the following remarks may be considered as a contribution to this important subject.

To enable me to present in a clear light the results I have reached, it becomes necessary to bring before you some theoretical points relating to the inversion of sugar solutions.

It is to Mitscherlich that we owe the observation that the deviating power of solutions of inverted sugar on polarised light varies with the temperature. It is, however, to Clerget that we owe a careful study of the subject. As far back as 1849 he published in the *Annales de Physique et Chimie* (vol. xxvi., 3rd series, p. 175) a process for rectifying errors in tests of commercial sugars by Soleil's saccharometer, by taking into account the deviation caused by a solution of inverted sugar, obtained by heating with hydrochloric acid a sugar solution, previously tested without inversion. In this communication, however, Clerget does not enter into theoretical considerations. He directs that sugar solutions shall be inverted by heating them with 10 per cent of their volume of concentrated hydrochloric acid up to a temperature of $68^\circ \text{C}.$,

* Read before the American Chemical Society, Feb. 6th, 1879.

taking about ten minutes in the operation. As soon as this temperature is reached, the solution is to be cooled down to some temperature between 10° and 35° C., as the table he gives for correction is calculated for temperatures between these limits, which, he says, "answer for all occasions which may present themselves in "Europe, as well as in the Colonies."

When Clerget proposed this new plan for making sugar analysis more correct, the subject of testing sugars was not a new one with him, for it is to him that we owe the idea of using Soleil's polarimeter as a special instrument for analysing sugar, and, as far back as 1845, he had published, in the *Bulletin de la Société d'Encouragement*, an account of Soleil's saccharometer. In his paper in the *Annales de Chimie et de Physique* he gives no theoretical reasons for the new process. For the theory of the correction of the direct test of the saccharometer, by taking into account the test after inversion, I am indebted to some formulas in Maudlebluh's Guide (*Leitfaden zur Untersuchung der Verschiedenen Zuckerarten*, Brunn, 1867, p. 56). This theory I will now proceed to lay before you.

You are aware that commercial sugars contain, besides pure cane sugar, a variety of substances, of which many exert deviating effects on a ray of polarised light. Among these is inverted sugar, which may be an immediate substance or a mixture of dextrose and lævulose. This latter supposition was advanced by Dubrunfaut, but has never been demonstrated in a satisfactory manner, although it would take volumes to collect what has been written about it and is being written about it every day.

The other bodies which accompany cane sugar have not been studied in such a way as to throw light on their nature, with the exception of aconitic acid, isolated by Dr. Behr, who gave an interesting account of his researches in a paper read before the Society two years ago.

A point of great importance in connection with the analysis of sugar by optical methods is that the impurities which are generally present are in such a condition that they seem to exert no effect on polarised light. The proof of this is to be found in the fact that with most sugars, particularly if the saccharometric test is above 90 per cent, the result, after the correction for inversion, is the same as given by the direct test. As many of the impurities reduce the alkaline solution of tartrate of copper, they are comprised under the head of *inactive glucose*, concerning the nature of which there are many unsatisfactory doctrines.

To explain how the direct test by the saccharometer may be corrected by subsequently inverting the solution on which this direct test was made, let us suppose that we have in the first instance a deviation, D . We may suppose that this deviation is the resultant of the following:—

- + C , deviation due to the cane sugar.
- − i " " " inverted sugar.
- + h " " " one or more substances, besides cane sugar, which turn the plane of polarisation to the right.
- − g , deviation due to one or more substances, besides inverted sugar, which turn the plane of polarisation to the left.

Then we may suppose that $C - i + h - g = D$.

In this equation the only known quantity is D , but what we want to know is C . To find this quantity we make use of this fact, that if the solution under examination is heated with 10 per cent of hydrochloric acid, the whole of the cane sugar will be converted into inverted sugar, while the other substances will not suffer any change. From some experiments which I made with artificial mixtures analogous, as far as I could judge, to those which constitute the impurities of commercial sugars, I am led to believe that by heating with concentrated acid a certain change takes place in the deviating power of the impurities of commercial sugars. This change is, however, very

slight, and as the impurities are in small quantities when compared to the cane sugar, the changes they suffer may be neglected, and we may suppose that, with the exception of the conversion of cane sugar into inverted sugar, no change takes place in the other constituents of the commercial sugar from the treatment with hydrochloric acid.

After this action of hydrochloric acid, aided by heat, has taken place, the solution is examined again in the saccharometer, and, after making a deduction for the volume of acid added, we will obtain a deviation, d . This deviation in ordinary sugars will take place on the negative side of the scale. We may suppose that it is the resultant of the same elements that made up deviation D , with the exception of the quantity C , now represented by the deviation due to the inversion of the cane sugar originally present. This quantity we will call $-I$, and we shall have $-I - i + h - g = d$. If now we wish to eliminate the quantities i , h , and g , we may easily do so by subtracting the second equation from the first, and we shall have $C + I = D - d$.

By a series of experiments Clerget established the relation between C and the corresponding value of I . This relation varies with the temperature, but we may suppose that the observation, after inversion, is taken at 28° C., at which a quantity of cane sugar which, before inversion, gives a deviation of 100 to the right, will show for the corresponding inverted sugar 30 to the left or -30 . At this temperature $C + I$ becomes $1.3 \times C = D - d$. As d is a negative quantity the algebraic subtraction is equivalent to an arithmetical addition, whence we draw the rule that at 28° C., the true quantity of sugar C is equal to the sum of the two deviations divided by 1.3, as—

$$C = \frac{D - d}{1.3}$$

This number 1.3 corresponds, as we have said, to 28° C. We may in the next place inquire: What are the corresponding numbers for other degrees of temperature? This brings us to the consideration of Clerget's table. For temperatures between 10° and 35° C., Clerget determined these numbers, and on consulting his table we find that for a quantity of cane sugar equal to 100 we must take for the quantity $D - d$ at 10° C., the sum 139, and that for every degree C. above 10° , up to 35° , the number representing the arithmetical sum of the deviations is equal to 139 minus one-half the difference between the number representing the temperature and 10° . Thus for 20, we have $20 - 10 = 10$ and $139 - \frac{10}{2} = 134$; for 28° , we have $28 - 10 = 18$ and $139 - \frac{18}{2} = 130$; for 35° , we have $35 - 10 = 25$, and $139 - \frac{25}{2} = 126.5$, &c.

If the same law should hold good below 10° , we must, to 139, add $\frac{1}{2}$ for every degree below 10° , and this leads us to establish that at 0° , the number corresponding to $D - d$ would be $139 + 5 = 144$. As this number 144 is one of great importance, allow me to recall to your attention that it means that if we have a solution of pure sugar which will produce a deviation of 100 divisions on the positive side of the saccharometer scale, if this solution is inverted by heating with 10 per cent of hydrochloric acid, this inverted solution, if tested at the temperature of 0° C., will, after making the correction for the quantity of acid added, produce a deviation of 44 on the negative side of the scale.

If now we suppose that the law which Clerget found for temperatures between 10° and 35° holds good for all other temperatures, we may easily obtain the deviation to the left of the same solution of inverted sugar at any temperature by subtracting from 44 one half the number expressing this temperature in degrees Centigrade. Thus we shall have for 10° , $d = -(44 - 5) = -39$; for 20° , $d = -(44 - 10) = -34$; for 28° , $d = -(44 - 18) = -30$, and for any temperature, t ,—

$$d = -\left(44 - \frac{t}{2}\right).$$

If now we draw a series of parallel equidistant lines, intersected by another series of lines, also parallel and equidistant, perpendicular to the first, we may take on the horizontal base line or line of the abscissas, a space between two lines corresponding to $x^{\circ}C.$, and we may suppose that the space between two lines of the other series corresponds to a division of the negative side of the saccharometer scale. If now, on every vertical line, starting from the base line, we take a length proportional to the numbers given by the equation—

$$d = -\left(44 - \frac{t}{2}\right),$$

and if we connect the extreme points of all these lengths by a line, we will find that it is a right line, because the equation—

$$d = -\left(44 - \frac{t}{2}\right) = -44 + \frac{t}{2}$$

is equivalent to the equation of the right line, $y = a + bx$, in which $d = y$, $a = -44$, $b = \frac{1}{2}$, and $x = t$.

If now we take again our equation,—

$$d = -44 + \frac{t}{2},$$

we will find that, if the law it expresses holds good to the end, the temperature at which the deviation becomes 0 is $88^{\circ}C.$

(To be continued.)

PROCEEDINGS OF SOCIETIES.

PHYSICAL SOCIETY.

Ordinary Meeting, May 10, 1879.

Prof. W. G. ADAMS, President, in the Chair.

New member:—Mr. J. Kestrell Evans.

Mr. WOLLASTON explained the construction of Gower's improved form of Bell's speaking telephone. The older form, made of wood or ebonite, is open to the objections that it has a very weak voice, soon gets out of adjustment from changes of temperature, and requires a twisted hand wire which is liable to break. Gower's form has a comparatively loud utterance, is constant, and does not require to be held in the hand, but may be laid on a table or hung on a wall, a speaking-tube leading from it to the operator's ear or mouth. The "call" for attracting attention is also within the Gower telephone itself; whereas, in the hand telephone it is an auxiliary apparatus. Every organ of the old telephone has been modified to form the Gower. The magnet in the Gower is of a horse-shoe form, very powerful, the two poles being brought very close together, and each pole is mounted with a small coil of fine wire. The diaphragm is much thicker and larger than the Bell diaphragm. The case is of brass, to expand equably, and a speaking-tube is fitted to the front of the diaphragm. The call consists of a musical reed attached to the diaphragm, so as to be opposite a small slit in the latter. To sound the call it is only necessary to send a sharp puff of wind up the speaking-tube, and the reed gives out a note which is heard throughout a room at the distant end. Speaking and cornet music was transmitted by the instrument exhibited, between the third storey over the hall and the meeting. It was very distinct and audible several feet from the receiver. Speaking done some thirty feet from the transmitter was also sent. Conversation was likewise carried on while considerable noise was being made in the room.

Prof. McLEOD remarked that the *timbre* of this telephone was very good.

Prof. W. F. BARRETT then gave an account of some attempts which he had made to overcome the induction

clamour on telephones caused by the ordinary telegraph currents on neighbouring wires. He had tried recently the Bell telephone on a line from Dublin to Armagh, 93 miles long, but the induction noises completely stifled the speaking, whereas the Edison transmitter gave good results. The clamour could be got rid of either by neutralising the induction currents or by eliminating the noises from the speech. He had taken the second line by experiment. Since the vocal currents differ from the induction ones in potential and period, he attempted to make the latter discharge across the line to earth by fine needle points, and from a heated spiral of wire, in a vacuum, leaving the vocal currents to pass on to the receiver, but without success. Also, since the vocal currents are alternately positive and negative, whereas the induction ones are of one sign, he tried to avail himself of the difference in discharging power of positive and negative currents, but without success. He then tried to take advantage of the difference of period or duration of the currents, the induction currents being longer. He therefore tried to break up the inductions by interposing a rapidly rotating current interrupter, and to make the sections of the musical note obtained interfere with each other by means of an acoustic interference-tube, but practically failed in this also. He mentioned these facts for the benefit of others who may be going over the same ground.

Mr. WOLLASTON pointed out that a perfect cure for induction on underground wires consisted in twisting the going and returning wire of the telephone circuit round each other.

Mr. WILSON then read a paper "*On the Divisibility of the Electric Light by Incandescence.*" By Joule's law the amount of heat developed in a circuit of resistance, R , by the passage of a current, $C = C^2R$; where R is the resistance of generators and connections, r , added to the resistance of the light emitter or incandescent wire, P . Therefore, since by Ohm's law,—

$$C = \frac{E}{R}$$

we have—

$$C^2 = \frac{E^2}{(r+P)^2},$$

and—

$$C^2P = \frac{E^2P}{(r+P)^2}.$$

From this equation the value of P may also be determined. C^2P is the amount of heat developed in the incandescent wire. He infers that the smaller the mass of the wire the higher the temperature generated in it, therefore the mass of the wire should be diminished until the fusing-point of the metal is almost attained. The question of divisibility resolves itself into our being able to divide a single incandescent source into a number of smaller ones giving the same total illumination. The author concludes that this can be done by arranging the sub-divided sources in "multiple arc," or parallel circuits, provided the total mass, length, and sectional area of the united sources be the same as in the original single source. The objection that increased radiation from the various sources would diminish the first total of light and heat can be met by making the smaller wires still smaller than is theoretically required so as to generate more heat. The author regards the "voltaic arc" as probably falling under the same law, the mass, however, being smaller in this case.

Dr. COFFIN then exhibited a Trouvé Polyscope, which consists of a small hand incandescent platinum wire electric light, designed for illuminating the more inaccessible cavities of the body in surgical examinations. The current is supplied by a Planté secondary battery, and the light is half enclosed in a small silver reflector, fitted with a convenient handle. The apparatus is portable. Dr. Coffin found that it was open to several objections, which he has remedied. First, the heat generated made the lamp so hot

that it could not be held to the body for more than a very short time. He overcame this by making the reflector of double silver plates, and circulating water between by means of india-rubber pipes from a bulb which can be worked by the patient himself, thus serving to distract his attention from the operation. Secondly, the secondary battery exhausts itself in twenty minutes, and the light therefore goes out, while from twelve to twenty-four hours are required to re-charge it. Dr. Coffin has superseded it by a Leclanché battery of 8 elements, made by Messrs. Coxetter and Sons, in which the carbon pole is replaced by a copper plate faced with platinum, and no porous diaphragm is employed. This gives a constant light for hours.

NOTICES OF BOOKS.

A Manual of Practical Chemistry: the Analysis of Foods and the Detection of Poisons. By ALEXANDER WYNTAR BLYTH, M.R.C.S., F.C.S., &c. London: C. Griffin and Co.

THE first portion of the title of this book, when standing alone as it does on the back, scarcely leads the reader to expect what is accurately enough described in the second part. The work consists of two somewhat distinct sections, the one treating of the analysis of foods, with especial reference to their impurities—intentional or accidental—while the other is devoted to the detection of poisons. Thus we have before us what might be pronounced two distinct treatises, whose bond of union is mainly due to the bookbinder.

At the same time we feel perfectly free to admit that each of these parts has been compiled in a judicious and conscientious manner. In dealing with adulterations the author has selected the best and most recent methods; he has appended certain useful legal decisions, and under each chapter he has given the bibliography of the subject. We are pleased to see that Mr. Blyth does not under-rate the difficulties of food analysis and of toxicology; he addresses himself to chemists, and does not, as some earlier writers have pretended to do, seek to make every man his own analyst.

In speaking of the analysis of milk we find that he, like most practical authorities, is able to confirm the substantial invariability of the "solids not fat" as existing in unsophisticated samples. He admits that the publication of a normal standard allows a certain amount of watering to be practised, but he fears that the bulk of commercial milk in this country is below the lowest estimate of the Society of Public Analysts. We must, moreover, warn him that it is not in all cases safe to discuss and criticise, even approvingly, published analytical processes. We have heard of one case where such criticism, which necessarily involves quotation, was met with the threat of an action for infringement of copyright! The remarks on unhealthy and abnormal milks are very interesting. The author considers it demonstrated that a disease similar to, if not identical with, tuberculosis may be propagated from animal to animal by means of the milk derived from a diseased cow. He considers that phthisis is very rare among cattle, and that when so affected their milk rapidly decreases in quantity. The chemical qualities of the milk secreted during cattle plague, pleuro-pneumonia, and anthracoid affections have not been duly ascertained.

The adulteration of tea Mr. Blyth thinks is substantially effected abroad, and is decreasing.

Turning to the toxicological section, we find a remark quoted from Dr. F. Mohr, which though mainly judicious is on one point open to question. This distinguished chemist says—"What relation sal-ammoniac, saltpetre, alum, tartaric, citric, and acetic acids, &c., have to toxicology we cannot conceive." Yet, unless we are greatly mis-

taken, a fatal case of poisoning is on record from an over dose of tartaric acid, taken by mistake for Epsom salts. Poisons are conveniently classed by the author under four heads—those which cause death almost immediately, irritants, irritant-narcotics, and poisons affecting the nervous system. The fourth group is again subdivided into narcotics, delirians, convulsives, and those producing complex nervous phenomena.

We cannot but consider it singular that though chrome is mentioned among the irritant poisons no instructions are given for its detection. This is the more to be regretted, as chromium compounds are used in the arts on a vast scale, and are present in the waste-waters from dye and colour works, &c., whence they may find their way into streams, wells, &c. Potassium bichromate, even applied externally, often produces unpleasant symptoms—a fact very familiar wherever "chrome-blacks" are much dyed.

In the section on animal poisons we find a most interesting account of the secretion of the cobra, which the author has carefully examined. The active principle is not a "germ" or organised ferment, but a well-defined and stable chemical compound, which Mr. Blyth has isolated and named provisionally cobric acid.

The whole work is full of useful practical information, and as far as we have been able to perceive it may be regarded as trustworthy. An exceedingly valuable feature—we believe original—is that the various pleas which may probably be raised on behalf of a supposed adulterator or poisoner are here pointed out, and the expert is thus forewarned. We have also very full instructions as to the various channels through which different poisons may be introduced into the human system.

A defect which we hope to see remedied in a future edition is the prevalence of typographical errors. The late Professor Gorup-Besanez is converted into Gorop Besaner; Crantz, the historian of Greenland, becomes "Crzaut," &c. In a tabular view of the composition of coffees from different localities, taken from a German source, some words are left untranslated. Thus we read of Jamaica and Ceylon "Plantagen." It should be "Plantations."

CORRESPONDENCE.

THE VITRIOL MANUFACTURE,

To the Editor of the Chemical News.

SIR,—Dr. Lunge seems to have misunderstood the purport of my letter. I did not intend to prove that an appreciable loss was actually experienced in the Glover tower; I wrote to point out that all fear of appreciable loss of nitre in the Glover tower had not yet vanished. I published some figures simply to show the possibility of such a loss being quite appreciable.

It would not from my point of view be necessary to reply to Dr. Lunge at all, since his remarks flow from this misunderstanding, but as some of these remarks are directed against my ability of weighing arguments, I feel it a duty to myself to make the following statements.

Until very recently the loss of nitre was by no means generally understood to be to so large an extent due to the reduction of nitrogen compounds. For example, Lock's Treatise on the Manufacture of Sulphuric Acid, dated 1879, decidedly inclines to the belief that Mr. Davis's views on the subject are the best explanations yet given of the loss of nitre. Dr. Lunge's treatise, just published in German, lays more stress on the loss of nitre incurred by reduction to nitrous oxide in the chambers. But the whole tenor of the explanations given of the reactions causing such loss is such as to produce the impression that these reactions occur only in isolated places in the chambers, viz., near the steam-jets, and is not at all calculated to give an idea of the magnitude of this loss, which Dr. Lunge now puts down at 75 per cent of the

nitre used. An attempt to give a measure of the loss even if only approximately, I have not found in this exhaustive and very able treatise. I consider it therefore a gain to have elicited from so high an authority as Dr. Lunge the admission (even if conditionally only) that the amount of nitre lost in a manner not yet clearly understood is 75 per cent, or more, of that used.

This "big margin" of course is affected by all the errors in estimating the losses from other sources, but it is quite certain that the statements I furnished err on the right side. Dr. Lunge differs from me only in the distribution of this "big margin." Whereas I think a portion of nitre (and I have many reasons for thinking so) is decomposed in the Glover tower, Dr. Lunge considers the whole of it destroyed in the chambers. He would have been more consistent if he had used a similar phrase to one occurring in his treatise, to the effect that certainly the Glover tower is in this respect no worse than any other denitrator.

To some extent, however, Dr. Lunge seems still to think that part of the "big margin" must be accounted for by the oxidation of arsenious acid to arsenic acid in the Gay-Lussac tower, when, he remarks, that my views conflict with "very careful investigations of M. Hjelldt." But permit me to point out that these "very careful investigations" are of the same type as Mr. Davis's. Both authors discovered arsenic acid in Gay-Lussac vitriol, both came to the conclusion that it was due to oxidation by nitrous anhydride at the expense of an equivalent of nitric oxide. Neither of the two gives experimental proof of this assertion; while Mr. Davis tries to find the resulting nitric oxide in the exit gas of a number of works, M. Hjelldt is satisfied with simply calculating the probable loss of nitre due to this supposed reaction. I have attempted in more ways than one to oxidise arsenious acid by means of nitrous acid in presence of strong sulphuric acid. I have allowed the reagents to act for seventy-two hours at ordinary temperature, and for several hours at temperatures up to 230° F., without being able to obtain a single bubble of any gas, either nitrogen, nitrous oxide, or nitric oxide. I have after repeated experiments come to the conclusion that arsenious acid cannot be oxidised to arsenic acid in presence of strong sulphuric acid so easily as to constitute a source of loss in the manufacture of sulphuric acid. Thus I am convinced that the "big margin" must be accounted for otherwise. Where the whole of the nitre is introduced into the Glover tower there remain only this apparatus and the chambers for the distribution of the loss.

Dr. Lunge thinks my fallacy consists in not having brought forward one particle of proof that any part of this "big margin" must be accredited to the Glover tower, and he produced, what is no doubt meant to be an excellent proof, that the whole of it must be ascribed to the chambers.

Dr. Lunge cites the results of two French works. In one of these (Scheurer-Kestner), where no Glover tower is used, 4.7 of nitre per 100 of sulphur were consumed, and Dr. Lunge calculates that 75 per cent were decomposed inside the chambers. If that is really so, how could the other works (Maletta, of Rouen) where nearly the same amount of nitre was used per 100 of sulphur, save 35 per cent of the nitre, simply by introducing a Glover tower. There would on Scheurer-Kestner's basis be at the outside 15 per cent to be saved, and based on Dr. Lunge's calculation I should say that the introduction of a Glover tower in Scheurer-Kestner's works would, as far as denitration is concerned, be practically useless. Dr. Lunge evidently has forgotten to charge Scheurer-Kestner's denitrator with part of the loss incurred. For a saving of 35 per cent in such works is only possible if the mode of denitration formerly in use is more disastrous than that by the Glover tower.

I have no experience of any other method of denitration except that by the Glover tower, and for ought I can say to the contrary it may be the very best form of deni-

trator, but that does not imply that the Glover tower is perfect, nor exclude the possibility of improvement.

In estimating the share of loss of the chambers I made use of Mr. Davis's figures. Mr. Davis defended these very figures as those he was most sure of. I have, however, some results of daily testings extending over three months of the exit gas of a set of chambers having neither Glover nor Gay-Lussac tower attached, and these results show that the amount of nitrogen escaping in form of its acids is within a small amount (20 per cent) equivalent to the nitre potted. All the conditions which are necessary to cause a reduction of the nitrogen compounds to a lower degree of oxidation than nitric oxide I conceive to be present in the Glover tower to an equal, if not a greater, degree than in the chambers. The sulphurous anhydride here in its greatest concentration is associated with all the steam evolved in the Glover tower, to react upon nitric oxide in the nascent state at a temperature much higher on the average than that of the chambers.

I can see no valid reason in what Dr. Lunge has brought forward to arrive at the conclusion that the loss in the Glover tower has "vanished into thin air," or, as he formerly expressed himself, is as small as a "differential quotient," whatever that expression is meant to convey. Still less can I see any great wisdom in so easily acquitting the Glover tower of all share in the destruction which is evidently going on in the system, and for which, I am fully convinced, the chambers are not wholly responsible. —I am, &c.,

FERDINAND HURTER.

Laboratory, Gaskell, Deacon, and Co.
Widnes, May, 1879.

THE VITRIOL MANUFACTURE.

To the Editor of the Chemical News.

SIR,—I do not wish to take up more of your space than I am obliged, but it seems to me necessary to state that in testing vitriol exits by a continuous system a Bunsen pump and meter is not necessary. An 8-litre bottle properly arranged is more than sufficient for eighteen hours' work, and if that quantity of gas be drawn out in such a manner as to accurately sample the whole number of cubic feet of gases passing, one cubic foot, or even less, will serve every purpose. One cubic foot of the exit gases in many of my experiments has been a sample of at least 1,200,000 cubic feet passing away. But what I principally want to point out (and which I did point out in my last, only it seems to have vanished in transit) is that both Hurter's and Lunge's theory of the chemical loss of nitre in the chambers will not work. Hurter ascribes 20 per cent loss, due to the action in the chambers. Lunge says 75 per cent. Now let us examine both of these.

A works with five large chambers was making 100 tons of vitriol per week, and they were using to produce this 3 tons of nitrate of soda per week; they used no towers, made always a good production from the ore burnt, and the escaping oxygen was 6.2 per cent by volume of the exit gases. The total acidity of the exit gases was always low—about 2.3 grains of Na_2CO_3 neutralised by one cubic foot. After working in this way for some years this firm decided to erect one denitrating column and two absorbing columns: directly they started to work there took place a great reduction in the quantity of nitre used, and the normal quantity soon became 16 cwts. per week.

Now let us examine these results closely, and the amount of nitre destroyed by the chambers must be reckoned on the original quantity used before the towers were erected, as your correspondents have not inferred that the introduction of towers will cause the destruction of nitre in the chambers to be any less, and we have no published experiments which even hint that this might be the case. These chambers, then, actually used 60 cwts. of nitrate of soda per week.

Hurter states twenty per cent is destroyed in the chambers; this is equal to 12 cwts.

Lunge states 75 per cent of the loss takes place in the chambers themselves; this is equal to 45 cwts.

Now the putting up of the towers can scarcely be thought to have exerted any influence upon the nitre-decomposing power of the chambers, as the drips and acids were worked to be of the same strength as before their erection. What was then the total nitrate of soda used weekly? 16 cwts., or 29 cwts. less than Lunge says the chambers alone will decompose. Lunge's theory, then, must evidently be a false one.

Hurter states that the chambers would decompose 12 cwts., leaving 4 cwts. to be accounted for in other ways. But he has accounted for them, and let us see how his theory works:—

Decomposed by chambers; 20 per cent of the	
60 cwts. used	12 cwts.
Withdrawn from system by exit gas; 10 per	
cent of the 26 cwts. used	1.6 "
Loss by imperfect denitration; 10 per cent of	
the 16 cwts. used	1.6 "
Loss from leakage; 5 per cent of the 16 cwts.	
used	0.8 "
Total loss	16.0 "
Total used	16.0 "

So that even on Dr. Hurter's own figures there is no nitre left for his theory of destruction in the towers.

I am glad that Dr. Lunge endorses my views with regard to the escape of nitric oxide. Many chemists have found it in company with oxygen, and if they will come forward and state their views I am willing to give the experiments which I have upon the subject.—I am, &c.,

GEORGE E. DAVIS.

Heston Chapel, near Stockport,
May 12, 1879.

PROBABLE BEHAVIOUR OF VARIOUS METALS AND THEIR COMPOUNDS WHEN SUBJECTED TO RAPID OXIDATION.

To the Editor of the Chemical News.

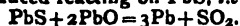
SIR,—Having received much assistance from chemists, and knowing the interest taken by them in the experiments which have been made at the works of Messrs. Charles Cammell and Co., Penistone, and at those of Messrs. John Brown and Co., Sheffield, I venture to hope that I may be favoured by the advice and assistance of some of your readers as to the probable behaviour of various metals and their compounds when subjected to rapid oxidation.

We will assume that the charges treated consist of sulphides, arsenides, &c., and contain a sufficient proportion of oxidisable materials to produce the necessary heat for their fusion, decomposition, and volatilisation of the sublimes. That iron, lead, gold, silver, copper, nickel, antimony, zinc, arsenic, and sulphur are present as well as sufficient silica to combine with the protoxide of iron produced, so as to form $2\text{FeO}, \text{SiO}_2$, and also an additional quantity of silica with the necessary lime or other bases to neutralise same so as to form a slag of specific gravity 3.4.

To avoid giving unnecessary trouble, I will briefly state my ideas as to their behaviour, and I trust your readers will kindly correct any errors I may make, as by so doing they will greatly oblige me, and thus assist me in my further experiments.

The Iron would be partially or entirely oxidised, and combine with the silica so as to form $2\text{FeO}, \text{SiO}_2$ + silicate of lime, &c.; part of the iron might remain in the regulus as FeS and Fe.

The Lead would probably distil off and be collected as PbS; but it is possible that metallic lead might be reduced by the PbO produced reacting on PbS, i.e.,—



If metallic lead is produced it would collect in the quiescent

hearth, and with it would be found the gold and silver, and probably some metallic copper.

The Nickel and Copper would be concentrated into a regulus, and would contain all the silver and gold, unless metallic copper or lead was reduced. If not too much oxidised the regulus would contain $\text{FeS} + \text{Fe}$, and if the oxidation was prolonged metallic copper would result, with which would be found all the silver, but I am not clear as to the gold. I should have supposed that the gold and silver would have gone together, but in an experiment at Penistone, July 17, 1878, a small quantity of metallic copper was produced, which Mr. Edward Riley found to contain 201 ozs. 4 dwts. 12 grs. of silver per ton, but only traces of gold, at the most 2 dwts., although the regulus from which it was produced contained 1 oz. 2 dwts. 20 grs. of gold per ton.

The Antimony and Zinc would be volatilised and collected as oxides.

The Arsenic would probably distil off as sulphide of arsenic, although part might be oxidised to arsenious acid.

The Sulphur not found in the regulus or combined with arsenic would partly distil off as sulphur, and the remainder would be oxidised and pass off as SO_2 .—I am, &c.,

JOHN HOLLOWAY.

Jeffrey's Square, St. Mary Axe,
London, E.C., May 10, 1879.

OBITUARY.

THOMAS WILLS, F.C.S.

MANY of our readers will have heard with regret of the very sudden death of Mr. Thomas Wills, who has acted as Secretary to the Chemical Section of the Society of Arts since it was first founded in 1874. From the *Journal* of the Society we learn that Mr. Wills was born in 1850, in Devonshire; he was educated at University College School and at King's College. In the early part of 1868 he became an assistant to Dr. Odling at St. Bartholomew's Hospital, and in the latter part of that year, on Dr. Odling being elected to the Fullerian professorship at the Royal Institution, Mr. Wills was appointed his official assistant. In 1873 he accepted the position of Demonstrator in Chemistry at the Royal Naval College. The subject to which he specially devoted himself was the application of chemistry to the manufacture of gas, and on questions connected with this subject he was rapidly becoming an authority. He held for some time the position of chemist to the Phoenix Gas Company. In 1873, previous to his official connection with the Society of Arts, he read a paper on "The Manufacture of Gas," and last year he gave a short course of lectures before the Society on "Explosions in Coal Mines." For several years he acted as Secretary to Section B of the British Association, and he was a member of the Association Committee for ascertaining the best methods of improving the illuminating power of coal gas. His most recent piece of work was in connection with the subject of electric lighting. Dr. Tyndall, in giving evidence upon the electric light before a Committee of the House of Commons, referred to Mr. Wills as having discovered that oxides of nitrogen were given off by the voltaic arc, thus rendering the light to that extent injurious. His death was most sudden and unexpected. He fell a victim to typhoid fever, which carried him off after an illness of less than a week. He was only twenty-eight years of age at the time of his death.

New Books.—The first volume of "A Treatise on the Manufacture of Sulphuric Acid and Alkalies," by Prof. Lunge, will, we understand, be published very shortly by Mr. Van Voorst. The work will be fully illustrated, and be complete in two volumes. The same publisher has in the press a supplement to Mr. C. Greville Williams's "Handbook of Chemical Manipulation."

MISCELLANEOUS.

South London School of Pharmacy.—The prizes for the Second B. Course were presented on Saturday, the 3rd inst., by the Secretary, Mr. W. Baxter, to the following successful competitors:—Senior Chemistry: Medal, Mr. Newbigin; Certificate, Mr. Lemmon. Junior Chemistry: Medal, Mr. Scammell; Certificate, Mr. Stevens. Botany: Medal, Mr. Betts; Certificate, Mr. Dutton. Materia Medica: Medal, Mr. Cook; Certificate, Mr. Stedman. Pharmacy and Practical Dispensing: Medal, Mr. Stevens; Certificate, Mr. Cook.

The Coal Tar Colours.—At the meeting of the Chemical Section of the Society of Arts, held on Thursday, the 8th inst., Mr. W. H. Perkin, F.R.S., read the first part of a paper on "The History of Alizarine and Allied Colouring Matters, and their Productions from Coal Tar." In this he gave a brief account of madder and its applications to calico printing, Turkey red dyeing, &c., and then proceeded to treat of alizarine and purpurin, the colouring matters of madder. He then gave a history of the discovery of their artificial production from anthracene, a hydro-carbon from coal tar, and the new processes by which it had been found possible to manufacture these bodies on the large scale, as well as the allied colouring matter, such as anthrapurpurin, &c. The subject was illustrated with experiments and specimens of the products referred to, as well as patterns dyed with the various colouring matters. The reading of the second part of the paper has been postponed from the 15th until the 22nd, when the history of the practical methods of carrying out this important manufacture will be dealt with.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 14, April 7, 1879.

Iodides of the Stannopropyls.—A. Cahours.—The author had formerly described a supposed iodide of sesquistanpropyl. He finds that it is a mixture of the diiodide of distannopropyl and of the iodide of tristannopropyl.

On Complementary Spinners.—E. Chevreul.—If a circle, one diametrical half of which is painted with any colour *a*, and the other half left white, is made to revolve at a speed between 60 and 160 turns per minute, the complementary colour of *a* appears on the white half. The author proposes these spinning disks as tests for colour blindness.

Focus of Heat Produced by Molecular Shocks.—W. Crookes. Presented by M. du Moncel.

Reply to M. Flammarion on the Declination of the Magnetic Needle.—Marié-Davy.—The author ascribes the anomalies pointed out by M. Flammarion to the methods employed in the calculation of the means in his tables.

The Gravi-volumeter.—A. Houzeau.—The gravi-volumeter is employed in a method of analysis which enables us to perform certain determinations not within the reach of ordinary volumetry. The author states that by its means he can determine with exactness and in less than twenty-five minutes the sulphates in a natural water, operating upon 10 c.c. without previous concentration. The instrument cannot be intelligibly described without the aid of the accompanying figure.

Detection of Fire-damp in the Atmosphere of Mines.—MM. Maillard and Le Chatelier.—The authors propose a lamp burning hydrogen gas, the feeble light of

which does not mask the blue flame produced by fire-damp within the gauze of safety-lamps. In this manner a proportion of hydrogen proto-carbide as low as 0.25 per cent is easily recognised.

Certain Conditions of Lactic Fermentation.—Ch. Richet.—The presence of oxygen facilitates the fermentation of milk. Up to 44° a rise of temperature intensifies fermentation. From that point up to 52° there is no modification, and above that point the fermentation slackens. Digestive juices augment the rapidity of lactic fermentation.

Amylaceous and Amyloid Granules of Eggs.—M. Dastre.—The author maintains that the amyloid bodies of M. Dareste are not starch and have not even its appearance. He remarks that micro-chemical reactions are not always trustworthy in cases of substances saturated with albumen and fat, and thus rendered impenetrable to reagents.

Determination of Sugar in the Blood.—M. d'Arsonval.—The author points out that the criticism put forward by M. Cazeneuve on methods employed by the late Claude Bernard have already been refuted. He states that greenish reductions betray either bad reagents or a defective manner of operating, and ought never to occur.

Methods Employed by Cl. Bernard in Determining Reductive Sugars in the Blood.—P. Picard.—The author admits that there exist in the organism, along with reductive sugars, other substances capable of acting upon Fehling's test, but he doubts if such can exist in the liquid obtained by the preliminary treatment adopted by Cl. Bernard? He considers that a saccharimetric examination of animal matter can lead to no accurate results as there are a great number of substances possessing a rotatory power.

Distribution of Phosphates in the Different Elements of the Blood.—L. Jolly.—The alkaline phosphates predominate in the serum, whilst phosphate of iron, though present in all the constituents of the blood, is especially accumulated in the globules.

The Formation of a Peculiar Amyloid Principle Peculiar to the Asci of Certain Pyrenomycetes.—L. Crié.—As a rule fungi possess no starch, but *Sphaeria Desmazieri* contains an amyloid mass formed in darkness by a protoplasm devoid of chlorophyll.

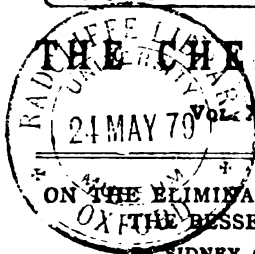
MEETINGS FOR THE WEEK.

- MONDAY, 19th.—Society of Arts, 8. "Recent Advances in Telegraphy," by W. H. Preece. (Cantor Lectures.)
Royal Institution, 3. "The Intellectual Movement of Germany from the Middle of the Last to the Middle of the Present Century," Prof. Hilkebrand.
- TUESDAY, 20th.—Civil Engineers, 8.
Royal Institution, 3. "Suggestions to Students and Readers of History," Prof. J. R. Seeley.
Society of Arts, 8. The Adjourned Discussion on Mr. W. Lloyd Wise's Paper on "The Government Patent Bill" will be resumed.
Zoological, 8.30.
- WEDNESDAY, 21st.—Society of Arts, 8. "Edison's New Telephone," Conrad W. Cooke.
Pharmaceutical, 8. (Anniversary.)
- THURSDAY, 22nd.—Royal Institution, 3. "Dissociation," by Prof. Dewar.
Society of Arts, 8. "The History of Alizarin and Allied Colouring Matters, and their Production from Coal-Tar," by W. H. Perkin, F.R.S.
Zoological, 4.
- FRIDAY, 23rd.—Royal Institution, 9. "Multiple Telegraphy, or Duplex and Quadruplex Telegraphy," Mr. W. H. Preece.
Society of Arts, 8. "The Harbour of Kurachee," W. J. Price, M.L.C.E.
- SATURDAY, 24th.—Royal Institution, 3. "On Swift," Prof. H. Morley.
Physical, 3.

TO CORRESPONDENTS.

ERRATUM.—Vol. XXXIX., p. 190, line 37, read "the former together with the H₂S set free," &c.

H. Allen.—The paragraphs were quoted from foreign journals, and it is impossible always to trace them to their proper source.



THE CHEMICAL NEWS.

Vol. XXXIX. No. 1017.

ON THE ELIMINATION OF PHOSPHORUS IN THE BESSEMER CONVERTER.*

By SIDNEY G. THOMAS, F.C.S., and
PERCY C. GILCHRIST, Associate Royal School of Mines, F.O.S.

THE non-removal of the phosphorus in the Bessemer converter, owing to which the great bulk not only of British, but of French, German, and Belgian ores, are still unavailable for steel-making, is a fact too familiar to metallurgists to need insisting on. The inquiry whether this unfortunate circumstance is due to causes absolutely inseparable from the conduct of the Bessemer process, or to others which are merely the accidents of a particular mode of constructing the apparatus, is obviously of vital importance. If the non-elimination be due to the intensity of the temperature, or to the short duration of the operation, or to both these causes combined, it is almost hopeless to expect that we shall ever be able to use ordinary unpurified pig-iron in the converter.

That it is to these essential accompaniments of the process that the phenomenon of the retention of phosphorus by Bessemer metal is to be ascribed is, it is believed, the generally received opinion, and one which has comparatively recently received the sanction of the weighty authority of such eminent metallurgists as Mr. Lowthian Bell, Dr. Wedding, Prof. Kerl, and M. Euverte.

An examination of the general condition attending the removal of phosphorus in puddling and refining operations, taken in connexion with the well-known action of silica on phosphate of iron at high temperatures, and the fact that in many other processes in which the temperature is very high the elimination of phosphorus is not apparently effected, seems, however, to justify the belief, which has doubtless suggested itself to other members of this Institute, that it is to the siliceous lining of the ordinary converter, and to the consequent necessarily siliceous character of the slag, that one defect of the Bessemer process is due. Under this conviction, at all events, experiments were commenced by the authors about three years ago on the effect of basic lining and basic additions in the several steel-making processes. Unfortunately the appliances at command were of a very imperfect character, and the results obtained, though highly encouraging, were, owing to defects in the miniature converter employed, which prevented our ever completely finishing a blow, not entirely conclusive as to commercially complete purification being possible.

While waiting the completion of an improved converter, which was unavoidably delayed for some time, we were encouraged by finding that M. Grüner, the distinguished professor at the Ecole des Mines, of Paris, in his "Treatise on Steel," published in 1867, laid great stress on the injurious influence of the siliceous character of the cinder and lining in the converter. M. Grüner, however, seems at that time to have regarded this as one only of three causes which prevent elimination of phosphorus, and proposes as a remedy the preliminary refining of phosphoretic pig before it is attempted to convert it.

With a new converter, a large number of experiments were made in the autumn of last year, which gave much more definite results. The lining used in these experiments consisted of limestone and silicate of soda, a mixture which had been found to answer well in earlier trials.

On laying some of the first of the results obtained from

* A Paper read before the Iron and Steel Institute. (Condensed.)

this six-pound converter before Mr. Martin, of Blaenavon, he at once recognised their importance, and from that time we have been deeply indebted to him for his unflinching and consistent support, and much valuable advice and assistance. The Blaenavon Company, also, without hesitation, undertook to put up apparatus to carry the experiments further, and has with great spirit fulfilled its promise to give every facility to test the value of the theory thoroughly.

In a vertical converter, taking from 3 cwts. to 4 cwts. of metal, results entirely confirmatory of those previously observed were obtained. In the six-pound converter liquid decarbonised iron could not be obtained; but in the new vertical converter this was readily done.

Some fifty or more blows were made in this vertical converter, and the products analysed; and it was found that, using a basic lining, it was generally necessary to continue the blow for over forty seconds after the flame dropped in order to bring the phosphorus down very low. With this proviso the elimination of phosphorus could be secured with absolute certainty. With a siliceous lining the retention of all the phosphorus in the metal was, as usual, equally invariable, even when, as in Mr. Bell's experiments, the blow was continued till a considerable proportion of the iron was oxidised. At the same time, more phosphorus and less silica would be found in the slag obtained under these conditions than appears to be the case when large quantities of metal are treated under similar circumstances.

Using a lining consisting of one part fire-clay and two of ganister, and a pig containing 1.44 per cent of phosphorus, the blown metal contained 1.63 per cent of phosphorus and the slag 32.5 per cent of silica, and 0.85 per cent of phosphorus. When, however, with the same lining, 40 lbs. of lime was placed in the converter before the pig was run in, though the lining wore away very much (as might be expected, there was a decided decrease of phosphorus in the blown metal, as shown below:—

P.	P.	P.	SiO ₂ .	CaO.
Pig used, 1.44	Blown metal, 1.23	Slag, 0.99	30.7	18.8
" —	" 1.07	" 1.81	31.0	25.1

It would seem that the presence of a considerable amount of lime in a not too siliceous slag is highly favourable, and on a large scale essential, to the removal of phosphorus. As it was manifest that phosphorus was not removed until the slag was sufficiently basic, the effect of large basic additions in combination with a basic lining was tried, with the object not only of obtaining a highly basic slag at an early stage of the blow but of rendering the operation independent of the wear of the lining by which alone the basic character of the slag is otherwise obtained and maintained. Advantage was taken of the fact that lime and oxide of iron are fusible in many proportions. The mixture generally used consisted roughly of one part by weight of "Blue Billy" and two of lime; this will melt in an iron crucible, and may be readily added in a molten condition. It was found that by throwing into the converter cheap basic materials consisting mainly of lime, even without previous heating, before the pig was introduced, very satisfactory results were obtained without over-blowing.

By using these basic additions, a large proportion of the phosphorus is eliminated while yet a considerable proportion of the carbon remains, a result which had otherwise only been obtained when there was a very considerable waste of lining.

With a 12-cwt. converter of the ordinary pattern, expressly put up by the Blaenavon Company, only a limited number of casts were made, owing to a deficiency of blast.

By the kindness of Mr. Menelaus, for whose kindly assistance we have to tender our sincerest thanks, we were enabled to try, at the old No. 3 pit at Dowlais, if the superior intensity of heat which might be expected from

the conversion of 5 or 6 tons of metal at a time affected the conclusions to which smaller experiments pointed. It was intended to line this converter with highly burnt basic bricks. The bricks intended for this purpose were, however, accidentally under-burnt, and so spoilt; hence recourse was had to a rammed lining of limestone and silicate of soda. This lining, which consisted of a siliceous limestone mixed with 9 per cent of a solution of silicate of soda, would, after the carbonic acid was driven off, contain nearly 20 per cent of silica. This, by greatly diminishing the effect of the ordinary wear of the lining in making the slag basic, rendered larger basic additions necessary than it was deemed prudent to make in the first two blows. In the first blow about 2 cwts. of lime were added cold, and in the second nearly 2 cwts. of lime and 1 of Bona ore. The slag was decidedly siliceous in both cases, and, of course, only a minute quantity of phosphorus was removed.

Blows.	Pig Used.		Blown Metal.		Steel.	
	I.	II.	I.	II.	I.	II.
Si ..	2'97	2'62	trace	trace	trace	trace
P ..	1'21	1'21	1'21	1'19 under	1'11	1'11
C ..	—	—	none	0'10	0'54	0'42
S ..	0'06	0'10	0'50	0'10	0'05	0'09

Mr. Jenkins, of Dowlais, has kindly furnished us with his analyses of the same blows. He finds 1'08 in the steel from the first blow, and 1'03 in the steel from the second blow.

The cinder with the steel contained in the first blow 38·8 per cent of silica, and in the second blow 36'8. The pig used was too high in silicon for phosphorus to be readily removed, without very large additions of lime.

In the third blow $3\frac{1}{2}$ cwts. of a mixture of two parts of lime to one of roll scale were thrown into the converter before the metal was run in. The metal was blown for eight or nine minutes and turned down (before the flame dropped). Nearly another $3\frac{1}{2}$ cwts. of the same mixture was then thrown in, and the vessel again turned up, when the flame dropped almost immediately. After being turned down for some fifty seconds, it was (at Mr. Martin's suggestion) again blown for nearly a minute. Though it appeared that the metal was overblown, the action, on adding spiegel, was not violent. A large skull was, however, left in the converter and ladle, and much slag was produced. The blow lasted ten and a half minutes. A rail made from one of the ingots deflected 98 inches with the blow of 1-ton ball falling 24 feet, the bearings being $3\frac{1}{2}$ feet apart. It was considered much too soft for rails.

The skull left in the converter was got out by blowing a charge of very siliceous non-phosphoretic pig. In the next (fifth) blow, 1 cwt. of a mixture of two of limestone and one of Elba ore was thrown cold into the converter before the metal was run in; rather over 3 cwts. of heated roll-scale was added subsequently, before the completion of the blow. During this blow the lining had to be patched at the breast.

The results appear to confirm the conclusion that, for the process to be of technical value, waste of lining and metal must be avoided by making large basic additions, so as to secure a highly basic calcareous slag at an early stage of the blow. In these trials, however, it was thought prudent to feel the way, and not add at once the very large amount of base which our theory demanded, the more so as we were not able to add the bases in the molten state. It is also made clear that a slag containing under 24 per cent of iron may be very effective in removing phosphorus.

After the five blows described the lining was found to be much worn, and not in a condition to admit of satisfactory repairs. Two tuyeres had to be renewed in the fourth blow; the rest stood well.

It is obvious that without a sufficiently durable, as well as refractory, basic lining, the simultaneous dephosphor-

isation and conversion of cheap pig in the Bessemer vessel cannot rank as a commercial process. Our early experiments rendered it clear that ordinary non-siliceous lime and limestone did not constitute by themselves a satisfactory lining material; nor were renewed trials, made after becoming acquainted with a patent dealing with their application, more successful. Magnesia, the use of which as a furnace lining has been suggested by M. Caron and others, is at once very expensive and—when used by itself—very tender. Nor did we, after repeated attempts, find it at all practicable on the large scale to use oxide of iron linings in any of the forms in which they have been used in puddling furnaces. After a very extended series of trials* it was, however, found that, by firing bricks made of an aluminosiliceous limestone at a very intense white heat, a hard and compact basic brick is formed. These bricks unfortunately labour under the defect of a liability to disintegration when exposed to the action of steam. By the use of certain aluminous magnesian limestones and equivalent combinations, and an otherwise similar mode of manufacture, this difficulty has been overcome. For certain purposes magnesian limestone, mixed with silicate of soda solution, forms an excellent material. To enter fully into the important subject of the precise conditions necessary for obtaining a satisfactory basic lining would exceed our limits, and the consideration of this, as of many other interesting points, must be reserved. The question of how far the heat due to the oxidation of phosphorus may replace that due to the combustion of silicon, the possibility of using in the converter low phosphoretic pig low in silicon, and the influence of silicon on the removal of phosphorus, are some of the subjects on which much remains to be said.

In advancing the proposition that the technical removal of phosphorus in the Bessemer converter is simply and entirely a question of cheaply producing a highly basic slag, containing under 20 per cent silica and over 30 per cent of lime and magnesia, and indicating the means by which this may be secured, we are not aware that we can shelter ourselves under any very distinct authority, though surmises as to the hypothetical advantages that might be expected, were the Bessemer slag less siliceous, have not been wanting. It is, however, only proper that we should remind the Institute that Mr. Snelus stated at its March meeting simultaneously with ourselves that he had removed phosphorus in a Bessemer converter lined with limestone. Of the circumstances of this experiment we are in ignorance. It is on the production of a basic earthy slag, by the addition of large quantities of calcareous bases, and without excessive waste of lining and metal, and the construction of a durable basic lining, that we venture to think the economic solution of the phosphorus problem depends.

It need hardly be said that the theory here advanced as to the practicability of commercially removing phosphorus in the converter extends, *mutatis mutandis*, to the Siemens and other open-hearth processes, where, in fact, many difficulties that are met with in the converter are absent. Dr. Siemens has indeed suggested the use of a lime lining in one of his furnaces, though he has since, with his customary candour, informed us that he has failed to devise means for its successful application. The present paper will have fulfilled its purpose if it induces metallurgists to reconsider the verdict, so fatal to the hopes of steel-makers, that "oxygen, whether in its free state or as oxide of iron, is almost entirely inert as regards phosphorus at the intense temperature which accompanies the Bessemer process."

In the nine months that have elapsed since our paper on the "Elimination of Phosphorus," &c., was prepared, the vigorous co-operation of Mr. Windsor Richards and

* For most valuable assistance in these, as at every period of this investigation, we are especially indebted to Mr. Thomas Griffiths, engineer at Blaenavon. We take the opportunity of at the same time thanking Mr. Gill, assistant in the Blaenavon laboratory, for his aid in the many hundreds of analyses made in the course of our inquiry.

Messrs. Bolckow Vaughan has enabled the authors to submit to the Institute further evidence in support of the views then put forward. That intensity of temperature is no obstacle to the removal of phosphorus, but, under proper conditions, highly favourable to this end, has been abundantly demonstrated by the results of the working at Middlesborough. It is indeed found that, other things being equal, the hotter the blow the better is the result. As regards the necessity for large additions of bases consisting mainly of lime, the experience afforded by some seventy or eighty operations is equally conclusive.

In continuation of some smaller experiments carried out in 1877 at Blaenavon, trials have been made of the effect of largely increasing the amount of oxide of iron added at the commencement of the blow, and diminishing the lime so as in a measure to assimilate the cylinder to that of a puddling furnace. As was anticipated, it was found that this could not be done successfully. In the first place, the loss from excessive boiling was very great; in the second, the phosphorus is only very imperfectly removed. A very considerable amount of ore may, however, be added after the greater part of the silicon is removed. The amount of bases which it is necessary to add with Cleveland pig generally exceeds considerably 2 cwts. per ton of pig treated, the exact amount being dependent on the wear of the bottom, and the percentage of silicon and phosphorus in the pig. The presence of an excess of earthy base in the slag seems an essential condition of success. The formation of a very fluid basic slag at an early stage of the operation is also of great importance, as it enables the phosphorus and carbon to be oxidised *pari passu*, or nearly so. It will be borne in mind that the basic addition has a double function; in the first place, to preserve the lining; in the second, to form a highly basic earthy slag, so as to afford a strong base with which the phosphoric acid may unite at the moment of its formation. On several occasions the experiment has been made of blowing in a basic brick lining without, or with very small, addition. The result is always excessive damage to the lining and a trifling removal of phosphorus. The density and compactness of the present lining material prevents it from playing the important part in the actual formation of the basic slag which was fulfilled by the softer and less durable linings first experimented on. The highly-fired magnesian lime bricks have, we venture to think, more than fulfilled all the expectations which were entertained as to their probable value.

It has also been ascertained that, providing a highly earthy basic slag is present, the removal of almost the last traces of phosphorus may be secured by continuing the blow for some time after the drop of the flame. The phosphorus in the presence of a strong base seems to protect the metal much in the same way as carbon and silicon, so long as there is present an excess of a strong base with which the phosphoric acid can unite at the moment of its formation. When, however, the phosphorus is reduced very low, the iron begins to oxidise as in an ordinary case of over-blow.

The question of the elimination of phosphorus being the primary matter with steel-makers, it is not proposed to do more than to mention that, under the circumstances which have been described as essential to the removal of phosphorus, a considerable proportion—viz., from 30 to 70 per cent—of the sulphur is also removed. Certain experiments made by the authors in conjunction with M. Ponsard, in November last, with a Ponsard furnace,—in the first of which over 80 per cent of the phosphorus and 50 per cent of the sulphur were removed under very unfavourable conditions,—quite confirm the opinion formed from smaller experiments, that the system described is as applicable to the open-hearth as to the Bessemer furnace. Those open-hearth furnaces which are provided with removal hearths offer peculiar facilities for the adoption of brick hearth or lining.

PEROXIDE OF HYDROGEN.*

By GEORGE E. DAVIS.

Most books tell us that H_2O_2 is made by a roundabout process with BaO_2 and HCl ; this might have been in days gone by, but now it is made principally by means of barium peroxide and hydrofluoric or fluosilicic acids. An insoluble fluoride of barium or fluosilicate is formed which separates at once and peroxide of hydrogen remains.

Now, if hydrofluoric or hydro-fluosilicic acids have been used, we need not expect to find hydrochloric, as this latter can only be present when impure fluoride is used, impure acids, or impure water. Of course it is absolutely necessary to examine carefully every sample of peroxide of hydrogen which may be purchased, for many statements have been made at various times respecting the purity of the bought article. One chemist assured me that the first lot he purchased was full of hydrochloric acid, whilst another chemist actually assured me that peroxide of hydrogen would not oxidise sulphide of sodium in vat liquors. More statements than these have been made respecting the efficiency of this reagent, but in each and every case it would have been more creditable to the author of the statement had he examined thoroughly the facts of the case before making complaint against one of the most useful articles used in chemical analysis.

Now, the first thing which requires to be estimated is the active oxygen, and this may be seen when I state that the sample which would not oxidise sulphide of sodium contained only a trace of active oxygen. Many methods may be devised for estimating the strength of this article. I have used the following three methods with satisfactory results:—

1 c.c. was put into Crum's tube over mercury and run underneath the tap; then 5 c.c. of a saturated solution of bichromate of potash, the whole gently agitated, the liberated oxygen measured and corrected for temperature and pressure. In working on a certain sample as a mean of several experiments, 1 c.c. gave 8.2 c.c. of oxygen at $0^\circ C.$ and 760 m.m.

By a second method 10 c.c. of the peroxide were boiled with 20 c.c. of sulphurous acid, and the sulphuric acid which was formed precipitated as barium salt and weighed; the sulphuric acid existing as such in the sulphurous acid being deducted from the precipitate. 10 c.c. oxidised 0.46 grm. SO_2 , or 7.08 grs.

By a third method 10 c.c. of the peroxide oxidised 146 c.c. $\frac{N}{10}$ protosulphate of iron. Now, if we calculate these we shall find that our first method gives the strength of the solution as 8.2 volumes, and they all, when expressed as in grammes of oxygen per litre, show:—

- a. 11.72
- b. 11.53
- c. 11.68

For general work the bichromate method is preferable, though when extreme accuracy is required the iron method may be employed if conducted in a stream of carbonic acid.

Now, as to the stability of peroxide of hydrogen. It has been stated quite recently, and by one of the members of this Club, that it is so unstable that simply drawing air through a solution of it will abstract the whole of its active oxygen. I don't know what amount of air is considered necessary, but I should say a great deal, and neither do I know what tests he used to ascertain that there was no peroxide left undecomposed. An alkaline solution of peroxide is partly decomposed by agitation only, but an acid or neutral solution is very stable. An alkaline solution of peroxide has been used over a very long time in connection with a continuous aspirator, 10 c.c. of peroxide and the necessary quantity of water

* A Paper read before the Faraday Club.

aspirated through at a speed of 15 cubic feet per twenty-four hours has shown an excess of peroxide at the finish of the experiment, even when caustic soda was present in excess also. I have evaporated a 10-volume solution of peroxide to half its bulk without driving off more than two-thirds of its active oxygen, but when reduced to that bulk caustic soda was added, the elimination of oxygen was so violent as to amount almost to explosion. When kept in solution in the laboratory a 10-volume solution of peroxide is moderately stable. During 1878 the following tests were made of a sample by the bichromate method over mercury:—

August 2	9'0
" 9	8'4
" 24	8'0
September 1	7'9
" 24	7'8
October 24	7'2

A few drops of ether were now added to 250 c.c. of that sample of October 24, and the following tests continued:—

November 7	7'2
" 24	7'2
December 1	7'2
" 24	7'2

Ether seems to preserve it, a fact which has been known for some time. This was also mentioned to me by Dr. Messel, of Silvertown, in August, 1878, a method I have used ever since. It is stated in all our text-books that peroxide of hydrogen is neutral to test-papers. Now, all commercial peroxide is faintly acid with the excess of hydrofluoric or hydrofluosilicic which is added and which should be estimated in each sample before using it.

I have used peroxide of hydrogen since 1873, and latterly in comparatively large quantities, and I have made it a rule to examine each purchase as follows:—100 c.c. of the peroxide was evaporated to dryness with 10 c.c. N soda, ignited, and taken up again with water. N acid was then added to neutrality, chromate of potash, and finally titrated with $\frac{N}{10}$ nitrate of silver. The results of many experiments upon this one sample showed that 100 c.c. of peroxide neutralised 0'2 c.c. of N soda and consumed 3'4 c.c. of $\frac{N}{10}$ nitrate of silver. On the evapora-

tion to dryness of the peroxide by itself a very pungent acid is liberated, and which can be easily told is not hydrochloric. In order to show the moderate stability of peroxide of hydrogen, I have lately had a sample sent me of commercial peroxide which was at least fifteen months old. When treated with bichromate over mercury it gave 7 volumes of oxygen, and I know that no special care, or indeed care of any kind, was bestowed upon the sample.

Having, then, a ready method of testing its contained amount of active material, it remains only for me to show that peroxide of hydrogen is an exceedingly useful oxidising agent, for it oxidises by reason of its loosely combined oxygen, and when an excess is added to any substance or to any solution, that excess is readily eliminated, leaving only as a residue that most neutral substance water.

There are many substances often seen in the laboratories of alkali works that cannot be readily and accurately examined except by its use; such as the total alkali and crude soda liquors or in black-ash, and certain qualities of soda-ash which contain sulphides or sulphites. In the testing of chamber exits it is extremely useful, inasmuch as sulphurous acid is not an acid easily titrated, for the normal sulphite of an alkali is not neutral to test-paper, and therefore on titrating a sulphite with a standard acid the point of neutrality is not clearly defined and distinct. This want of clearness in the ending has been stated by some chemists to be due to the carbonic and nitrous acids

present in chamber exits, but add peroxide of hydrogen to the solution and all difficulty vanishes, the sulphurous acid is oxidised to sulphuric, which acid is easily titrated.

I will now proceed to give examples of its use in various methods of analysis. Some samples of soda-ash contain so much sulphite of soda that it is impossible to estimate the amount of alkali accurately by means of standard acid. As an example of this kind of ash I give the following analysis of a sample of ash made from caustic salts, which contained by the acid test in the ordinary way 30 per cent of alkali without peroxide of hydrogen, whilst the addition of a few c.c. brought down the actual percentage to 21. The escaping carbonic acid carries away a great deal of sulphurous acid, but it does not do this if peroxide of hydrogen is present.

Sodium sulphide	0'098
" sulphite	22'982
" sulphate	2'207
" chloride	33'048
" silicate	1'214
" carbonate	18'704
" hydrate	12'800
Water	8'747

99'800

Peroxide of hydrogen is very useful in the analysis of black-ash; a few c.c. added to the liquor under examination gives at once the total alkali, which number needs no correction for sulphides, hyposulphites, &c.

This reagent is very useful for many laboratory oxidations, such as the oxidation of iron salts, colouring matters, experiments on bleaching, &c., &c. In fact the applications of this useful substance are legion. I first commenced to use this reagent for testing vitriol exits in 1873; but its application then was very limited as I had to make my own, and a perfectly pure peroxide of barium is not a very easy thing to make. In those days peroxide of hydrogen was looked upon as a sort of *rara avis*, to be found only in the laboratories of schools of chemistry, and its use was not much extended until a pure peroxide of hydrogen entered the market as a commercial article, to be bought and sold in the same way as other reagents.

Heaton Chapel, May 12, 1879.

EXPLOSIONS IN FLOUR-MILLS AND COAL-MINES.

By WATSON SMITH, F.C.S., F.I.C.

I BELIEVE it was about the year 1871 that the great explosion took place at the Tradeston Flour Mills, near Glasgow, but not having any particulars with me here, I cannot speak with absolute certainty as to exact date. As this was a catastrophe attended with loss of life and great damage to property, a searching investigation was made into its causes. Several theories were propounded, but none appeared to be satisfactory or to meet the circumstances but the one which I was enabled to give. This explanation was suggested to me on reading a short note in *Dingler's Polytechnisches Journal*, of which an abstract appeared under the head of "Technical Chemistry" in the *Journal of the Chemical Society*. The author, who wrote from Austro-Hungary I believe, had noticed how artificial lightning was made in theatres by blowing lycopodium seed through a flame, and he found that dry flour would also do for the purpose. Then he argued that flour dust in quantity might also be caused to flash off, given a due admixture of air and the introduction of a spark or flame. In this way he sought to account for sundry explosions in flour-mills in his part of the world. The sparks might be struck by the stones coming into slight collision, or incautious approach with a naked flame might fire the

mixed atmosphere of the stove-room in a flour-mill. The explanation I gave, which appeared in the *Glasgow Herald*, was based on this, and, so far as I know, I think I may claim priority as to the bringing forward of this view, as far as either Great Britain or America are concerned; at least I understood the Scottish Royal Society's Commission of Inquiry on the subject gave me at the time the credit of it. The drift of my explanation was the following:—"If the grinding-stones be supposed to get slightly out of gear at any time, as they quite possibly may, they may come in contact with each other so as to strike sparks amongst the fine flour-dust suspended cloud-like in the atmosphere immediately in the neighbourhood of the stones. Such an attenuated combustible mixture as flour-dust suspended in air approaches the condition of a mixed gas, and, the proportions being favourable, an explosive mixture. The whole atmosphere of the stove-room is often laden with this dust. Now, imagine a single grain of flour-dust inflamed by contact with a spark from the stones, or by a naked flame brought into the room, what is to hinder this inflammation from passing instantly, and like a flash, from grain to grain, the atmospheric oxygen supplying a common means of combustion? quite analogously to the case of an explosive gaseous mixture. Carbonic acid and steam are thus generated suddenly, and at an elevated temperature, and the sudden expansion in a closed space, like the interior of a mill, produces the pressure resulting in disastrous explosive effects."

Berthelot has calculated that the sudden increase of volume which may take place is amply sufficient to account for explosions. Under certain normal condition, in which great rapidity in the revolution of the stones is attained, and the grain is unusually dry, one can even, with Mr. L. Smith, conceive that sufficient heat may be developed to cause ignition, especially if some hard foreign body—such as a piece of wood, for instance—by any accident should get in with the grain in grinding. Its attrition and pulverisation might, perhaps, cause ignition of the dust produced. Still, an abnormal rapidity of revolution would also, in case of slight collisions of the stones, cause showers of sparks to fly, or an ordinary lantern or bare flame might be brought into the room, and these would be quite sufficient to inflame the atmosphere, given favourable proportions of flour-dust and air existent at the time. This it seems to me is an explanation nearer at hand, and also more probable than the other.

Mr. Langbeck in his paper (*CHEMICAL NEWS*, vol. xxxix., p. 191), though thinking himself justified "in opposing these explanations and substituting another one," neither, in point of fact, does practically oppose them, nor does he offer anything in substitution but an unlikely surmise—a mere "smell of impure hydrogen," on which opinions again might differ. This surmise he tries to support by experiments, which, however, on the whole, go more to second the explanations already given above than his own. The experiment with the coal-gas, as recorded, is abortive to prove anything on the subject; and, finally, his opposition to the explanation given, and his substitution of another one is a complete fiasco, for he winds up asking the way to the source of his own surmise. Having attained this, but failing to find his way out of it again, he puts it to the reader to find the way out for him in a couple of questions further. And this is opposing an explanation and finding a substitute! To both questions I for one would reply "Certainly, No!" as far as the case in point is concerned. If one of them were admissible it might surely be expected that explosions should occasionally be heard of in our bakers' ovens. Perhaps Mr. Langbeck is not aware that a well authenticated explosion is on record caused by the falling of the contents of a flour-sack. The cloud of flour-dust came in contact with a gas-flame, and a very considerable explosion occurred. The *Journal of the Chemical Society* of London for this month contains a brief account of Berthelot's views on the subject.

It is now a pretty well ascertained fact that some

otherwise slight coal-mine explosions may be expanded into great disasters, owing to the clouds of coal-dust raised into and diffused through the atmosphere of the workings; such an atmosphere, especially in presence of traces of fire-damp, being thus suddenly converted into an explosive mixture. This view, too, explains the frequently observed fact that an explosion in a coal-mine often sweeps over very considerable tracts of the workings. I am told that Prof. Marreco, of Newcastle, has investigated and experimented on this subject.

It has occurred to me that in view of these similar causes of, and liability to, explosions, both in flour-mills and in coal-mines, the electric lamp would be the best and safest means of illumination that could be employed. Such a light could be confined in a glass globe, hermetically sealed, and would be a fixture of course. Means of signalling could also be adjusted in coal-mines. Thus most stringent and rigorous measures would be rendered feasible to prevent the introduction of matches, lights, or lamps and lanterns of any kind into the workings. At the present cost of the electric lamp it might also with advantage be used in the stove-room of flour-mills, and others where clouds of flour-dust may be raised at any time. This recommendation may of course have been already made; if so I have not observed it, and on this ground may plead excuse for failing to acknowledge prior claims.

Zürich, May 9, 1879.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, May 15, 1879.

Dr. WARREN DE LA RUE, President, in the Chair.

THE minutes of the previous meeting were read and confirmed. The following certificates were read for the first time:—C. J. Wilson, G. S. V. Wills, W. H. Kerr, G. R. Tweedie.

THE PRESIDENT then called on Mr. WAREINGTON to read a paper "On Nitrification (Part II.)." A. Müller was the first to advance the opinion (*Chem. Soc. Journ.*, 1873, 1267) that nitrification is due to the action of a ferment. Schloësing and Müntz proved this to be the case (see Part I. of the author's paper), and from recent experiments show that soils which induce nitrification have this power destroyed by exposure for one hour to 100° C., and that ordinary moulds and mycodermis injure rather than promote nitrification. The author also mentioned that the above experimenters were at present engaged in an attempt to isolate and cultivate the organism, which promised good results. The objects of the author were to ascertain the influence of light, temperature, variations in the composition, and concentration of the solutions on the process of nitrification, the rate at which it progresses, and the relation of the nitric acid produced to the ammonia consumed. In nearly every case exposure to light prevents nitrification, and in all cases the exposure hindered the process to a marked extent. The mould which develops in a solution containing tartrates is incapable of effecting nitrification. The presence of carbonate of calcium seems to be indispensable to the growth of the ferment. The author pointed out the significance of this fact as explaining the absence of nitrites and nitrates in soft peaty waters, and as bearing on the utility of applying lime, &c., to peaty soils rich in nitrogen, in a form unfavourable for absorption by plants. A very small amount of organic carbon is requisite. An extensive series of quantitative experiments is given as to the effect of temperature. The upper limit of temperature at which nitrification takes place has not been determined. 40° C. is,

however, fatal to the process, which can proceed at 10°, and probably at still lower temperatures. In all cases there is a period after the addition of the ferment during which no appreciable effect is produced. This period the author terms the period of incubation. This period is considerably shortened by increasing the temperature. Thus, in a solution containing 640 milligrams. AmCl per litre the period was at 10° 78 days; at 30°, 19 days. As the solutions become stronger the period increases. Thus, a solution containing 80 milligrams. the period was only 31 days at 10°, or 12 at 30°, instead of as above 78° and 19°. The presence of bacteria does not promote nitrification. The author discusses the interesting question why in some cases nitrites, and in others nitrates, are produced. When the ammonia disappears before the nitrous acid is converted into nitric acid, the nitrites left in solution are very stable: if, however, the oxidation of nitrites sets in before the ammonia has been consumed nitrates are formed with great rapidity. In no case is the whole of the ammonia obtained as nitric acid. Some experiments were made as to the part played by the heating effect of the sun's rays in preventing nitrification, as a temperature of 40° proved to be fatal. It was found that a solution sheltered partially from the heat, by a screen of alum solution, but fully exposed to the light, of the sun, nitrified sooner than a similar solution exposed to the heat and light. In both cases nitrites were formed which were very permanent. The paper concludes with some interesting experiments on the conversion of nitrites into nitrates by the ferment. This change apparently takes place only in the dark, and a ferment which is quite competent to convert ammonia salts into nitrites is apparently not necessarily competent to convert the nitrites into nitrates. Some solutions, however, which are nitrifying seem to possess this power in a high degree. The progress of nitrification is not uniform; it begins slowly, increases in rapidity, and after reaching a maximum again diminishes. The strongest solutions nitrified by the author contained 180 parts of nitrogen per million.

The PRESIDENT said the author had investigated the subject with great acumen. It was interesting to observe that the process might produce either nitrites or nitrates.

Dr. WRIGHT then read a paper "On the Alkaloids of the *Veratrum* Family (Part III.)," by C. R. A. WRIGHT and A. P. LUFF. Alkaloids of *Veratrum album*.—The authors have examined the alkaloids extracted from 12 kilos. of dried roots by percolating with alcohol acidified by tartaric acid (1 part per 200 of roots), evaporating to a small bulk, addition of water, filtration from resin, and treating with a slight excess of caustic soda and ether. After repeated washing with ether an insoluble precipitate was left, which seemed to consist principally of a base hitherto undescribed, which the authors name pseudo-jervine, $C_{29}H_{43}NO_7$. It is snow-white, and melts at 299°, crystallising anhydrous from alcohol; with sulphuric acid it gives a yellow solution, gradually turning green. The ethereal solution contains, besides small quantities of pseudo-jervine, several other alkaloids, which can be separated by shaking the crude ethereal solution with aqueous tartaric acid, and treating the mixed tartrates with soda and a smaller bulk of ether: a residue is left, containing pseudo-jervine, an amorphous alkaloid named by the authors Veratralbine, and Jervine. Jervine, $C_{26}H_{37}NO_3$, forms a sulphate almost insoluble in hot and cold water. It crystallises with two molecules of water, melts at 239°, and gives with sulphuric acid the same colours as pseudo-jervine; the sulphate of pseudo-jervine is, however, tolerably soluble in water. The second ethereal solution deposits on spontaneous evaporation crystals of jervine mixed with another base, which forms a readily soluble sulphate. This base gives with sulphuric acid a red colouration, hence the authors suggest the name Rubijervine. It melts at 237°, and resembles in many respects pseudo-jervine; forms with crystallised salts, and crystallises anhydrous as $C_{26}H_{43}NO_2$. The ethereal mother-liquor of these crystals dries up to a varnish consisting

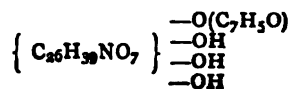
chiefly of veratralbine, $C_{28}H_{43}NO_5$. A small quantity of another base is present, yielding veratric acid on saponification. The mixture of veratralbine and this base is powerfully sternutatory, but this property is lost by boiling with alcoholic potash. Hence it is probable that the sternutatory constituent is veratrine (Couerbe). Neither jervine, pseudo-jervine, rubi-jervine, nor veratralbine excite sneezing. Veratralbine gives with sulphuric acid a red colouration resembling that given by cevadin and veratrine. No evidence of saponification or other decompositions was obtained on boiling these bases with alcoholic potash, the minute quantity of supposed veratrine excepted.

Dr. WRIGHT then read a paper "On the Alkaloids of the *Veratrum* (Part IV.)." Alkaloids of *Veratrum viride*.—On treating about 18 kilos. of dried roots precisely as described in the foregoing paper, the first treatment with ether left undissolved some pseudo-jervine, the tartrates obtained from the ethereal solution yielded no veratralbine, but jervine crystallised out from the second ethereal solution on standing. Traces of rubijervine were observed. The ethereal mother-liquors dried up to a powerfully sternutatory amorphous mass, closely resembling the "veratralbine" similarly obtained from *V. album* roots. It gave on analysis, however, $C_{32}H_{49}NO_9$, the formula of cevadin, and on saponification it yielded about the theoretical quantity of cevadic acid with a trace of veratric acid. The following table represents the approximate yield of the different bases from the two roots per kilo.:-

	<i>V. album</i> .	<i>V. viride</i> .
Jervine	1.30	0.20
Pseudojervine	0.40	0.15
Rubijervine	0.25	0.02
Veratralbine	2.20	trace
Veratrine	9.05	less than 0.004
Cevadine	apparently absent	0.43
	4.20	0.80

The jervine and pseudo-jervine from *V. viride* agreed in melting-point, properties, analytical numbers, &c., with the specimens obtained from *V. Album*.

Dr. WRIGHT then read a third paper "On the Alkaloids of the Aconites (Part IV.)." by C. R. A. WRIGHT and A. P. LUFF. Japanese aconite roots.—The authors have examined two different batches of roots, treating them with alcohol acidulated with tartaric acid, evaporating, adding water, making alkaline with sodium carbonate, and then shaking with ether. Repeated treatments with ether failed to dissolve all the alkaloid present, some being obstinately retained by the alkaline fluid; this appeared to be ac-crystalline. The ethereal extract, after purification by shaking with aqueous tartaric acid and treatment of the crude tartrate solution with soda and fresh ether, gave by spontaneous evaporation crops of crystals. These crops were fractionated and re-crystallised; all gave numbers indicating the formula $C_{56}H_{88}N_2O_{21}$. Treatment with hot concentrated tartaric acid failed to produce any change in the analytical numbers, whence the authors conclude that the substance is not a mixture of two bases, $C_{33}H_{45}NO_{11}$ and $C_{33}H_{43}NO_{10}$. The authors have named this base Japaconitine. It melts at 185° to 186°, and closely resembles aconitine. On saponification it splits up into benzoic acid and a new base, Japaconin. Japaconin closely resembles aconin, but on treatment with benzoic anhydride it forms a tetrabenzoylated instead of a dibenzoylated derivative. A tetrabenzoylated body is also formed by heating japaconitine with benzoic anhydride, aconitine giving a dibenzoylated body. The authors adopt the following view as to the constitution of japaconitine, admitting the existence of a base—



$$\begin{array}{c} (\text{C}_{26}\text{H}_{39}\text{NO}_2) - \text{O} \text{C}_7\text{H}_5\text{O} \\ | \qquad \qquad \qquad // \qquad \qquad \qquad \text{O} \\ | \qquad \qquad \qquad \text{O} \\ | \qquad \qquad \qquad // \qquad \qquad \qquad \text{O} \\ \text{C}_{26}\text{H}_{39}\text{NO}_2 - \text{O} \text{C}_7\text{H}_5\text{O} \end{array}$$
$$(\text{C}_{26}\text{H}_{39}\text{NO}_7) \begin{array}{c} \diagup \text{OH} \\ = \text{O} \\ \diagdown \text{OH} \end{array} .$$

Dr. WRIGHT, in reply, said that he was not convinced as to the identity or otherwise of his alkaloid and the one obtained by Paul and Kingzett. From the data given by the above authors he had stated that the properties, &c., of their alkaloid agreed with those of pseudoacotin, and he still held to this statement. He had only withdrawn it as far as his own alkaloid was concerned, which was certainly not pseudoacotin. He did not dispute the accuracy of nitrogen determinations by volume, but if only a per

The SECRETARY then read a "*Preliminary Note on some Reactions of the Ammonio-chloride of Magnesium known as Magnesia Mixture*," by H. D'ANCY POWER. The author has observed that most potassium and some sodium salts precipitate magnesium hydrate from a solution of ammonio-chloride of magnesium. Potassium iodide possesses this property in a marked degree. Thus the addition of 15 c.c. of a 10 per cent solution of potassium iodide with 10 c.c. of ammonia to 5 c.c. of magnesia mixture (prepared by dissolving 5 grms. of magnesium oxide in 40 c.c. of hydrochloric acid, and then adding 60 c.c. of ammonia and filtering), after standing twenty-four hours gave a precipitate which, when washed and ignited, weighed 0.046 grm.; it was pure MgO, so that 46 per cent of the total MgO was precipitated. Potassium bromide gave under similar circumstances a precipitate weighing 0.002 grm. Further results are promised.

The two following papers were taken as read:—

"*The Composition of Cows' Milk in Health and Disease*," by A. WYNTER BLYTH. The results of this research are the separation of two alkaloidal bodies as normal constituents of milk; the separation of a substance, probably a glucoside, derived from plants, &c., eaten by the cow; a quantitative estimation of the different constituents of milk; analyses of samples of milk derived from cattle in an unhealthy state. The separation of the milk alkaloids:—A litre of milk is divided into three equal parts, to one of which a litre of water is added, the casein is precipitated in a flocculent condition by the cautious addition of acetic acid, and, finally, by passing carbonic acid, a clear yellow whey is obtained, which is separated by decantation and filtration and used to precipitate the second portion; the whey from this is similarly used to precipitate the third portion of milk. The yellow whey is boiled and filtered to get rid of albumen, and to the filtrate an excess of the solution of nitrate of mercury used for urea estimation is added. The precipitate which falls contains the two alkaloids, any albumen, and urea as mercury compounds. It is washed and decomposed with sulphuretted hydrogen, &c. The first alkaloid, which the author proposes to call Galactine, is thrown down by acetate of lead; the lead salt has the composition $(\text{PbO})_{23}\text{C}_{54}\text{H}_{78}\text{N}_4\text{O}_{45}$. Galactine is a white, brittle, neutral, tasteless, non-crystalline mass, soluble in water, insoluble in alcohol; it is precipitated by Sonnenschein's and Scheibler's reagents. Excess of lead used to precipitate the galactine is removed and nitrate of mercury added, which throws down an alkaloidal colouring matter. Lactochrome, the empirical formula of the mercury salt, $\text{HgOC}_6\text{H}_{18}\text{NO}_6$. Lactochrome is a bright red-orange resinous body, softening at 100° , soluble in water and hot alcohol. In addition to these alkaloids the author has separated two substances, CH_3O_3 and $\text{C}_3\text{H}_3\text{O}_4$, reducing copper solution, which he regards as decomposition products of one substance, and as derived from food eaten by the cow. They are obtained by precipitation with ammonia and tannin, after separating the above alkaloids. The author gives the following as the average composition of healthy cow's milk:—Milk-fat, 3.50 per cent (oleine, 1.477; stearin and palmitin, 1.75; butyrin, 0.270; caproin, caprylin, and rutin, 0.003); casein, 3.93; albumen, 0.77; milk-sugar, 4.00; galactine, 0.17; lactochrome, not determined; bitter principle, 0.01; urea, trace; ash, 0.70 (K_2O , 0.1228; Na_2O , 0.0868; CaO , 0.1608; Fe_2O_3 , 0.0005; P_2O_5 , 0.1922; Cl , 0.1146; MgO , 0.0243); water, 86.87. As regards milk from diseased cows the author concludes that a cow suffering from even very acute disease may give milk differing in no essential feature from normal milk, whilst local affections of the udder may often be easily recognised. Analyses of milk from cows suffering from mammitis, pneumonia, phthisis, &c., are given.

"*Notes on the Effect of Alcohol on Saliva and on the Chemistry of Digestion*," by W. H. WATSON. The author finds that ptyalin is more rapidly or effectually precipitated from simple aqueous solutions than from saliva. The separation is aided by heating to 100°F . 18 grs. of absolute alcohol added to 200 grs. of saliva and 10 grs. of starch produced in an hour less extractive matter by about one-quarter than a similar mixture containing no alcohol; a similar reduction in the quantity of sugar produced was also effected by the addition of alcohol. The author made some experiments as to the effect of slightly acidulating the mixture of starch and saliva with hydrochloric acid. He arrived at the following conclusions:—By the addition of a small amount of acid the action of the saliva is decidedly increased, while the retarding influence of alcohol is not lessened by the presence of the acid. He also points out the bearing of these experiments on the process of digestion in the stomach, where the starchy matters and saliva are mixed with the gastric juice.

During the reading of the last three papers Dr. GILBERT took the Chair.

The Society adjourned to June 5, when the following papers will be read:—"On Gardenip," by Dr. Stenhouse and Mr. Groves; "On the Theory of the Fractional Distillation," by F. D. Brown; "On the Action of Organo-zinc Compounds on Quinones," by F. R. Japp; "On Chlorotannic Acid," by J. W. Mallet; "On Indigo-purpurin and Indirubin," by E. Schunck; "Third Report to the Chemical Society on some Points in Chemical Dynamics," by Dr. Wright and Messrs. Luff and Reanie.

CORRESPONDENCE

DETERMINATION OF NITRIC AND NITROUS ACID.

To the Editor of the Chemical News.

SIR,—In the *CHEMICAL NEWS*, vol. xxxix., p. 205, Mr. Davis quotes a remark of mine on Crum's mercury method (which Mr. Davis calls *his* method, although it was published many years ago, and used by several chemists previously), as if I had not rendered justice to him. I had certainly pointed out that the correction for temperature and pressure is required for something less than "extreme accuracy," as Mr. Davis has put it, considering that it amounts sometimes to more than 10 per cent, and I do not see that I can retract that. I have lately published tables which allow these corrections to be made by a simple reading off, without any calculation (*Dingler's Journal*, vol. cxxxi., p. 522).

With reference to Mr. Warington's paper, read before the Chemical Society at its last meeting, I beg to point out that in the *Berliner Berichte*, vol. xi., p. 439, I have described a number of tests, proving the accuracy of Crum's mercury method for nitrous and nitric acid, for a mixture of both, and for mixtures with arsenious acid and with glucose. In the just-quoted paper in *Dingler's Journal* I have described some precautions required for correctly working the process with my "Nitrometer."

Although Mr. Davis cannot in any sense claim the process as his, yet I have to thank him for bringing it forward again, as I had entirely overlooked it before, and I believe alkali works' chemists were generally in the same situation, and are equally indebted to Mr. Davis.

Would you kindly allow me on this occasion to correct an omission of a few words in my letter, printed in your issue of the 2nd inst., which obscures the meaning of the passage. On p. 194, col. 2, line 5, it should stand—"The total loss of nitre, *apart from that in the chambers themselves*." That this is the meaning of the passage is proved by the sequel.—I am, &c.,

GEORGE LUNGE.

Technical Laboratory of the Federal
Polytechnic Schools, Zürich.

PREVENTION OF ESCAPE OF SULPHUR-GASES DURING THE CHARGING OF PYRITES FURNACES.

To the Editor of the Chemical News.

SIR,—Having erected a set of new furnaces and acid plant seven years ago in the neighbourhood of a town where some of the local authorities were hostile to the manufacture, it was therefore of the greatest importance to prevent any escape of sulphur-gases. I was led to study the best means of preventing the escape during the time of charging. I hit upon a very simple and efficient plan, which had the recommendation of costing nothing. It was simply to close all the ash-pit doors before the door of the furnace to be charged was opened. The necessary draught being forced to enter through the only open door entirely prevents the escape of gas, and the workman

has only to open the ash-pit doors again the requisite distance to admit the proper amount of draught to each furnace. This plan I found to have the advantage of making the fires burn more regularly, as the fireman, from having to open and close the ash-pit doors oftener, had his attention directed to them, and so the draught was better regulated.—I am, &c.,

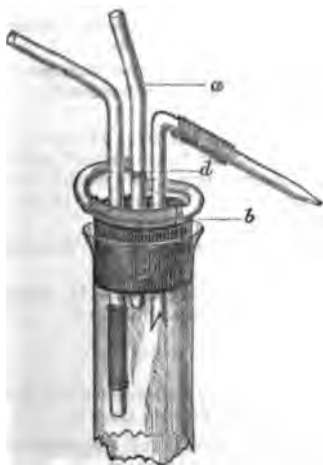
E. H.

May 20, 1879.
PS.—I may say that the furnaces and ash-pits were furnished with closely-fitting doors.

WASH-BOTTLES WITH CONTINUOUS JET.

To the Editor of the Chemical News.

SIR,—In the *CHEMICAL NEWS*, vol. xxxix., p. 19, Mr. Foye called attention to a wash-bottle giving a continuous jet of water, and Mr. Mallet, in your issue of May 2, has described a more complete but complicated form of the same. In both these cases the essential feature is a piece of caoutchouc tubing, so arranged as to permit the air inside the bottle being cut off from the external air by pressure with the finger. Will you kindly permit me to say that I described a bottle having this arrangement, and also the Bunsen valve, in a paper read before the Chemical Society in March, 1877. A short abstract appeared in your journal. The following is an illustrated description of the wash-bottle:—



It has two tubes fused into it at right angles, which are bent round upon the top of the cork, and connected by a piece of good caoutchouc tubing (*b*). The tube (*a*) is stopped between the two junctions of the side tubes by a piece of cork (*d*), or by being fused up. When the operator wishes to use the bottle he holds it as usual, placing his forefinger upon the caoutchouc (*b*), so as to prevent air passing through it, and blows through the mouthpiece into the bottle. After this the jet of water continues to issue with almost undiminished force for about three-quarters of a minute, while it may be stopped at any time by releasing the pressure upon the caoutchouc tube (*b*). Water may be poured out through the new tube (*a*) as usual. It is convenient to bend the lower end of the tube dipping into the water; this enables the operator to blow out the last portions of water. The advantages of the bottle are as follows:—The jet will issue for some time after the operator has blown into the bottle, and can be directed upwards into a beaker by inclining the bottle. The jet can be stopped at pleasure. The water passing through the new tube when the bottle is inverted can be regulated to a drop by pressing upon the rubber. The mouthpiece never gets hot, nor does gas or steam pass through it into the mouth. When the water is boiling, if the bottle be occasionally

gently shaken, enough steam is liberated to impel the jet. Lastly, the bottle can be used as an ordinary one by blowing into the new tube without pressing the caoutchouc. After the tube (*a*) is made or purchased there is no extra trouble in making the bottle. A caoutchouc stopper is the best to use, but a good cork answers perfectly well. Messrs. Jackson, of the Barbican, at my request have kindly consented to keep a few of these tubes in stock. Several of these bottles have been in constant use for two or three years in the Chemical Laboratory of the Royal College of Science, Dublin, and have given great satisfaction.—I am, &c.,

THOMAS BAYLEY.

14, Mincing Lane, London,
May 5, 1879.

THE VITRIOL MANUFACTURE,

To the Editor of the Chemical News.

SIR,—In the *CHEMICAL NEWS* (vol. xxxix., p. 205) Mr. G. E. Davis has referred to a statement I made to him regarding the loss of nitre. What I said then—I was speaking from memory—was perhaps scarcely justified by the facts; still, the quantity of nitre unaccounted for in our case is very much less than that mentioned by Dr. Hurter. The quantity of nitre used during the six months was 1·66 per cent on the stone burnt, equal to 37·18 lbs. to the ton. If we assume the real nitrate of soda present to be equal to 95 per cent, this would be equal to 35·31 lbs. There was found in the outlets nitrous gases equivalent to 15 lbs. nitrate of soda for each ton of stone burnt, which would give 42·5 per cent on the quantity used, or four times that mentioned by Dr. Hurter. If we add to this 10 per cent for imperfect denitration, and 5 per cent for mechanical losses, we get a total of 57·5 per cent unaccounted for. During the six months these trials were made much acid was drawn from the chambers for manure making, and this acid always contained a quantity of nitre; indeed, so much was this the case on some occasions as to interfere with the workmen while mixing in the vat. It is impossible to say how much was lost in this way, but it will be readily seen that much more than 45 per cent of the nitre is thus accounted for, without putting down 20 per cent as reduced to nitrous oxide, as is done by Dr. Hurter. I may say that we are still finding in the outlets between 40 and 50 per cent of the nitre used. But is it not probable that the different results obtained are due in a great measure to the different way in which the chambers are worked? This we know from experience—that where the chambers are worked with the acid strongly impregnated with nitrous gases much less nitre is required than where the acid in the chambers stinks of sulphurous acid.

Much has been said regarding the state of oxidation of the nitrous gases found in the outlets, but no one seems to have made any direct attempt to ascertain what that state of oxidation is. Directly after Mr. Davis published his observations on the reducing action of As_2O_3 in the absorber, I commenced a series of experiments to settle the question, at least as far as these works were concerned. In all fifty-five tests were made, and out of these in forty-three cases the gases existed as N_2O_3 and N_2O_4 , while in the other twelve the gases existed as N_2O_2 and N_2O_3 . It was necessary in making these experiments to assume that the nitrous gases were present in only three forms of oxidation, viz., as N_2O_2 , N_2O_3 , and N_2O_4 , and that only two of these, either $N_2O_2 + N_2O_3$ or $N_2O_3 + N_2O_4$, were present at the same time. This assumption is, I think, warranted by the properties of the gases, for we should scarcely expect N_2O_2 and N_2O_4 to exist together side by side without reacting upon each other.

Mr. Davis has attributed the presence of N_2O_2 in the outlets to the reducing action of As_2O_3 exerted while the acid is passing down the absorber. Now I have reason to believe that under certain conditions sulphurous acid is competent to reduce the nitrous acids. I give below two

instances, copied from my note-book, which will tend to support this belief:—"The chambers had been working with very great regularity for a length of time, the quantity of nitre used being equal to about 1 per cent on the stone burnt, and the quantity found in the outlets equal to about 12 lbs. on the ton of stone. On the 16th the chambers were looking badly, and I examined the pipe between the absorber and the last chamber when the gases smelt quite sharp. Accordingly, I expected to find a large quantity of SO_2 present (in the test solutions) this morning. I was surprised to find only a trace as usual, while that of nitre had more than trebled, viz., 43·31 lbs. to the ton of stone."

The next case is still more conclusive:—"Two days ago the acid from the absorber contained nitrous acids equivalent to 2·45 per cent of nitre; this morning it contained 1·07 per cent. During this time, the Glover tower being out of repair, has not been working, and it was thought advisable to put the same acid over and over again through the absorber. While this was going on the chambers have been stinking, and it was inferred that a large quantity of SO_2 would be found going away. This has not proved to be the case, as only a trace has been found in the test-bath yesterday and to-day, although 6 cubic feet of the gases have been tested on each occasion. The nitre found in the outlets has been very large, viz., 102·8 lbs. per ton of stone yesterday and 120·7 lbs. per ton of stone to-day."

Now these two cases would point to the fact that SO_2 is capable of reducing the N_2O_3 , in solution in the acid, to N_2O_2 ; thus, $\text{N}_2\text{O}_3 + \text{SO}_2 = \text{N}_2\text{O}_2 + \text{SO}_3$, and that this reaction may take place in the absorber.—I am, &c.,

E. JACKSON.

Laboratory of Messrs. T. Adkins and Co.,
Smethwick, near Birmingham.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 15, April 14, 1879.

Anomaly in Magnetic Observations at Paris.—C. Flammarion.—The author contends that the explanations of these anomalies put forward by M. Marié-Davy are not satisfactory.

Bulletin de la Société Chimique de Paris,
No. 5, March 5, 1879.

Recent Paper by M. B. Aronheim.—P. Schützenberger.—M. Aronheim having stated in the *Berichte der Deutsch. Chem. Gesell.* that on attempting to prepare the acetates of chlorine and iodine he had obtained mere mixtures of acetic and hypochlorous acids, or of chloride of iodine and iodic acid, M. Schützenberger refers to numerous chemists who have seen and examined, or have themselves prepared, the products in question.

Chloro-stannates of the Rare Earths.—P. T. Cléve.—The composition of the lanthanic, ceric, didymic, and yttric chloro-stannates is quite analogous to that of the corresponding chloro-platinates. The author has verified the researches of M. Marignac on the earths of the gadolinite. No doubt can exist concerning terbia. He has obtained a small quantity of ytterbia, but his researches are not completed.

Russian Chemical Society, December 7/19, 1878.—M. Beketoff communicated a paper on the determination of the atomic heat of hydrogen in its combination with platinum. M. Andrianowsky has examined the action of

aluminic chloride upon acetic and sulphuric anhydrid MM. Riordinine and Saytzeff describe diallyl-isopropyl carbinol as a colourless liquid, of sp. gr. 0·8512, boiling-point 182° to 185° . MM. Semianitzine and Saytzeff have investigated the oxy-valerianic acid obtained the oxidation of allyl-dimethyl-carbinol. MM. Beilstein and Kourbatoff have obtained nitro-phthalic acid oxidising nitro-naphthalin by means of chromic anhydrid

Researches on Sulphates.—A. Etard.—The sulphates of the magnesian series, or of the alum series, are from representing all the combinations which sulphuric acid forms with the metals. Many of these salts may be represented by the general formula $\text{M}_2(\text{SO}_4)_6\text{N}_2$. He examined the double sesqui-sulphate of iron and alum (acid); that of iron and chrome (acid); the corresponding neutral salt, &c. The sesquisulphate of aluminium and manganese is a powder of a beautiful sky-blue.

Relations between the Chemical Composition and the Mechanical Properties of Steels.—V. Deshay.—Carbon renders steels rigid and elastic, increasing the elastic tension, but their resistance to rupture diminishes if 0·500 is exceeded. Manganese renders steels rigid and elastic and increases their elastic tension, but the elongation and contraction remain considerable, which give good resistance to a shock. Silicon plays the same part as carbon, rendering steels hard, and slightly diminishes elongation. Sulphur decreases the breaking strain; the resistance to a shock. Phosphorus renders steels deficient in body, and, if its proportion exceeds 0·250 per cent, fragile on receiving a shock. Chrome adds to manganese but more energetically.

Thermic Formation of the Compounds of Carbonic Oxide with the other Elements.—M. Berthelot.—Already noticed.

Specific Heats and Melting Heat of Gallium.—Berthelot.—The specific heat of liquid gallium is 0·08 that of solid gallium 0·079. The melting heat is +19°. The specific atomic heat in the liquid state is 5·59; in solid 5·52.

Reciprocal Displacements of Weak Acids.—Berthelot.

Rotatory Power of Styrolen.—M. Berthelot.—The two papers have been already noticed.

Justus Liebig's Annalen der Chemie,
Band 195, Heft 3.

Products of the Action of Caustic Potassa upon Sulpho-mesitylenate of Potassium.—Oscar Jacobs.—An extensive paper in which the results obtained by Fittig, Hoogewerff, and others are critically examined. Among the products of the reaction are mesitol, oxy-mesitylenic acid, oxy-trimesinic acid, and oxy-uvitic acid. These compounds with a number of their derivatives are described at length.

Analysis of Organic Bodies Containing Halogens or Nitrogen.—Hugo Schiff.—The author describes with slight modifications the method proposed by Piria thirty years ago. The substance is weighed in a small platinum crucible, and in case of chlorine or bromine fills up with mixture of 1 part anhydrous carbonate of soda and 4·5 parts of lime, the whole being thoroughly mixed together by means of a platinum wire. The whole is then covered with a larger platinum crucible and the pair are inverted so that the bottom of the larger crucible is turned downwards. The space between the two crucibles is filled with the same mixture of lime and soda, the lid is put on, and the whole heated to redness over a strong Bunsen burner. The reaction is completed in about 10 minutes. In case of an iodine compound carbonate of soda is used without lime. The arrangements for the determination of nitrogen cannot be intelligibly described without the aid of the accompanying plate.

The Molecular Magnitude of Indigo.—Dr. E. v. Sommaruga.—The author finds the vapour-density of indigo = 9.45, and concludes that the formula for indigo must be $C_{16}H_{10}N_2O_2$ and not the half of these numbers.

Communications from the Laboratory of Applied Chemistry of the University of Erlangen.—These communications embrace the following papers:—Detection of Ethyl-diacetic Acid in Urine, by A. Hilger; on Solanin and its Decomposition Products, by A. Hilger; and, Certain New Salts of Uranyl, by Dr. R. Sendtner.

Influence of the Isomerism of the Alcohols and Acids on the Formation of Compound Ethers.—N. Menschutkin.—An important paper which does not admit of useful abstraction.

Contribution to a Knowledge of Polyporic Acid.—C. Stahlschmidt.—The author obtains hydropolyporic acid by treatment with caustic alkali, examines the salts of the new acid, obtains nitropolyporic acid by treating polyporic acid with concentrated nitric acid and certain chlorine compounds by the action of potassic chlorate and hydrochloric acid.

Contributions to a Knowledge of Ammoniacal Mercury Compounds.—H. Gerresheim.—An examination of Millon's base. The author announces an intention of shortly making known an application of this compound in water-analysis.

Moniteur Scientifique, Quasneville.
April, 1879.

Researches Conducted in the Laboratory of MM. P. Monnet and Co. at La Plaine.—P. Monnet, F. Reverdin, and E. Noelting.

Presence of Meta-nitro-toluol in Nitro-toluol.—The authors, in experimenting upon the acetylic derivative of a commercial toluoydin free from aniline, were led to suppose that in addition to the ortho- and para-derivatives it contained a little of the third isomer or meta-compound. To decide this question they have studied the nitro-toluol used in the manufacture of the toluoydin in question, and found their suspicions confirmed.

The Part of Meta-toluydin in the Manufacture of Aniline-Red.—The function of ortho- and of para-toluydin in the preparation of magenta has been studied by M. Rosenstiehl and other chemists; it seemed to the authors interesting to complete these observations by oxidising meta-toluydin either alone or mixed with aniline and the two other toluydins, especially as the presence of this base in a small extent had been traced in commercial toluoydin. They have heated various mixtures containing meta-toluydin with arsenic acid under the usual conditions for the manufacture of magenta and have extracted the colouring matter produced, and tested it by dyeing comparative samples:—

Base or Mixture of Bases in Equal Molecules.	Shade of the Product of Oxidation.
1. Aniline	Violet (violanilin).
2. Ortho-toluydin	Red.
3. Para-toluydin	Yellowish brown.
4. Meta-toluydin	Brown.
5. Aniline and ortho-toluydin	Red.
6. Aniline and para-toluydin	Red (para-rosanilin).
7. Aniline and meta-toluydin	Violet.
8. Aniline, ortho- and meta-toluydin	Yellowish and greyish red.
9. Aniline, para- and meta-toluydin	Red, slightly violet, and greyish.
10. Aniline, ortho- and para-toluydin	Red (ortho-para-rosanilin).
11. Ortho- and para-toluydin	Red.
12. Ortho- and meta-toluydin	Red, violet, and greyish.
13. Para- and meta-toluydin	Brown.
14. Ortho-, para-, and meta-toluydin	Red, violet, and greyish.

Artificial Alizarin.—C. Græbe and C. Liebermann.—The first part of this bulky treatise deals with the chemistry of artificial alizarin as understood down to 1874. Of the second part, which gives an account of subsequent improvements as far as the end of 1877, we shall shortly give an abridgment.

Contributions to the History of Benzol-green and Malachite-green.—This paper consists of certain memoirs by O. Dœbner and by O. and E. Fischer, taken from the *Berichte der Deutschen Chemischen Gesellschaft*.

Industrial Society of Mulhouse.—Proceedings of the Sessions of the Chemical Committee, December 11, 1878, and January 15, 1879.—Nothing is here described which has not been already noticed. The particulars of M. Kopp's process for the spectral analysis of dye-ware are not yet made known.

On Amber.—M. Helm.—Amber in entire fragments is permeable to water. It contains as much as 4 per cent of sulphur in the state of organic combination. This sulphur has probably been absorbed by the fossil resin in the state of hydrogen sulphide subsequent to its formation. The author describes another fossil resin, gedanite, which differs from amber by containing a smaller proportion of oxygen, and is softer, more fusible, and more soluble in ether. It is free from succinic acid.

Verhandlungen des Vereins zur Beförderung des Gewerbflusses. February, 1879.

The Manufacture of Magenta by Coupier's Process.—Dr. C. Häussermann.—The author remarks that Coupier's process has hitherto been adopted in but few establishments, and although the magenta thus produced is often preferred, especially as a raw material for the preparation of superior blues, it is not yet finally decided which of the two methods is preferable. For Coupier's process it is essential to use an aniline oil of sp. gr. 1.006 to 1.007 at 15°. It is converted into hydrochlorate with the presence of as little water as possible. To prevent the aniline hydrochlorate when dehydrated by a temperature of 140° from adhering too firmly to the sides of the vessel, two-thirds of the aniline to be used is slightly supersaturated with hydrochloric acid, concentrated till the temperature rises to 140°, and the remaining one-third of the aniline is then poured into the melted cooling mass. This operation is conducted in enamelled pans provided with a condensation apparatus. The mixture of aniline and aniline hydrochlorate is then introduced into the melting pans, adding to 100 parts of aniline about 50 parts of pure nitro-benzol, and by degrees 3 to 5 parts of iron filings.

March, 1879.

Ammonia Soda and the Production of Ammonia in Coke-burning.—Dr. Frank.—The author points out that the weak point of the Solvay process is its dependence on the supply and consequent price of ammoniacal salts. He insists upon the importance of utilising the volatile products—ammonia, tar, &c.—from coke works, which he considers would effect a revolution in all trades based upon the destructive distillation of coal.

Test for Mercurial Vapours.—At a recent meeting of the Society of the Physical and Natural Sciences of Bourdeaux M. Merget recommended paper steeped in the ammoniacal solution of nitrate of silver, or in chloride of palladium, as reagents for mercurial vapours much more sensitive than gold foil. This test-paper is very sensitive; a slip of sheet-copper plunged into a liquid containing 1 part of mercury in 10,000 remained bright after immersion, but if exposed to the ammoniacal nitrate of silver paper it occasioned a characteristic black spot. He finds that even when solidified mercury emits vapours in appreciable quantity.

MEETINGS FOR THE WEEK.

- SATURDAY, 24th.**—Physical, 3. "On a New Harmonograph," W. J. Wilson. "On a New Induction Balance," Prof. Hughes.
- MONDAY, 26th.**—Royal Institution, 3. "The Intellectual Movement of Germany from the Middle of the Last to the Middle of the Present Century," Prof. Hillebrand.
- Royal Geographical, 1. (Anniversary.)
- TUESDAY, 27th.**—Civil Engineers, 8.
- Royal Institution, 3. "Suggestions to Students and Readers of History," Prof. J. K. Seeley.
- Anthropological, 8.
- Society of Arts, 8. "The Contact of Civilisation and Barbarism in Africa, Past and Present," by Edward Hutchinson.
- WEDNESDAY, 28th.**—Society of Arts, 8.
- Geological, 8.
- THURSDAY, 29th.**—Royal Institution, 3. "Dissociation," by Prof Dewar.
- Royal, 8.30.
- Philosophical Club, 6.30.
- FRIDAY, 30th.**—Royal Institution, 9. "Colour-sense in Insects," Mr. Grant Allen.
- SATURDAY, 31st.**—Royal Institution, 3. "On Swift," Prof. H. Morley

TO BE SOLD BY AUCTION pursuant to an Order of the High Court of Justice Chancery Division made in the Cause of Hutchinson v. Norwood 1878 H. No. 14 with the approbation of the Vice-Chancellor Sir Richard Malins the Judge to whose Court the Cause is attached by Mr. Thomas Colclough Leete (of the firm of Messrs. Branch and Leete) the person appointed by the said Judge at the Law Association Rooms Cooke Street Liverpool on Thursday the 29th day of May next at 3 o'clock in the afternoon.

The Valuable and Extensive Chemical Works situate at Widnes in the County of Lancaster known as Messrs. John Hutchinson and Co.'s No. 1 and 2 Works established by the late Mr. John Hutchinson and carried on by his executors since his death. The property comprises a superficial area of 71,385 square yards of land of which 16,859 square yards are freehold and 54,526 square yards are leasehold principally held for long terms of years at low ground rents 221½ square yards of the leasehold land have been sublet and chief rents amounting to £30 6s. 7d. will be included in the sale.

The situation of the works is exceptionally favourable for communication both by Railway and water they are divided by the St. Helen's Canal which affords a convenient means of access on peculiarly advantageous terms and they immediately adjoin the River Mersey fronting which there is a pierhead in connection with No. 2 Works where goods can be loaded and discharged free from toll.

The Works have Railway communication with each other and are also immediately connected by means of sidings with the London and North Western the Sheffield and Midland and Great Northern Railways and with the Widnes Dock in which as well as in the Canal there are rights of berthage connected with the works. They also possess ample facilities for drainage and for the disposal of alkali waste on favourable terms.

The buildings plant machinery and apparatus are on a very extensive scale the greater part being of the most approved construction they are now in first rate condition and are capable of producing weekly 500 tons of salt-cake 225 tons soda-ash 75 tons caustic soda 30 tons soda crystals 25 tons bicarbonate soda and 20 tons recovered sulphur for all of which productions these works bear a most favoured character in the market. There is also plant almost ready for working designed for the manufacture of about 50 tons per week of bleaching-powder by Deacon's process.

No. 1 Works consist of the necessary buildings machinery and plant for the manufacture of vitriol salt-cake soda-ash soda-crystals bicarbonate soda and sulphur with Smith's filters and joiner's workshops a well yielding weekly about 500,000 gallons Railway weighing machine Railway sidings and tramways throughout and loading and discharging berths on the Canal and Widnes Dock.

No. 2 Works consist of the necessary buildings machinery and plant for the manufacture of vitriol salt-cake soda-ash caustic soda bleaching-powder and sulphur with Offices and Lavatory Coach-house Stabling and Out-buildings Cooperage and Cooperage Store Locomotive Shed Smith's filters and joiner's workshops and railway wagon repairing shop railway weighing machine saw-mill mortar-mill and engine a well (weekly capacity 500,000 gallons) railways and tramways throughout and a pierhead and wharfage on the River Mersey.

The Works will be sold in One Lot as a Going Concern and the purchaser will be required to take the stock stores implements and utensils at a valuation.

The premises may be viewed on application to Major Cross at Widnes and particulars and conditions of sale may be had of Messrs. Gregory Rowcliffe and Rawle Solicitors 1 Bedford Row Messrs. G. S. and H. Brandon Solicitors 15 Essex Street Strand London of Messrs. Part Woodcock and Co. Solicitors Wigan and of Messrs. Branch and Leete Auctioneers Hanover Street Liverpool.

SOAPERY.—To be Let, on advantageous terms, the ALBERT SOAP-WORKS, Sheffield; the only Soapery situated in the midst of a very large and increasing population. It contains five pans—one to cleanse 9 tons, two 6 tons, one 4 tons, and one 3 tons; Ley Vats; Hot Room; Steam-Engine; Circular-saw Room; and all suitable apparatus. Tenant may enter on very favourable terms, and at a moderate rental.—May be seen working at any time, by applying at 69, Broad Street, Sheffield.

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THE CHEMICAL NEWS.

VOL. XXXIX. No. 1018.

ON ELECTRIC DISCHARGES IN ATTENUATED GASES, AND ON CERTAIN PHENOMENA IN GEISSLER'S TUBES.*

By H. EUGEN GOLDSTEIN.

IN H. Goldstein's memoir, communicated to the Berlin Royal Academy of Sciences, May 4, 1876, we find the following results:—

The existence of a twofold conduction in attenuated gases, hitherto assumed, and corresponding to the so-called positive and negative light, cannot be retained. With the exception of the peculiar stratification of the negative (cathodic) light the author has succeeded in imparting all its supposed distinctive properties to the positive light.

If the calibre of the tube widens in the direction of the current, the discharge appears as a positive light. Under reversed conditions it is negative. In a number of experiments the cathodic discharge behaves as if taking place through a number of fine pores. By means of great rarefaction or, by increasing the intensity of the discharge, H. Goldstein succeeded in converting the spectrum of the positive light in tubes filled with air, nitrogen, or hydrogen into a spectrum of the cathode light. The phenomena observed by Reitlinger and Kuhn (*Pogg. Ann.*, 141) are explained by the fact that these experimentalists made use of a tube whose gaseous contents were absorbed by the electrodes during the passage of the current. The author has repeatedly met with such tubes. Under the influence of a magnet positive light behaves exactly like the negative, and is even affected by a smaller magnetic force. If the anode is in an axial position it is surrounded by a luminous cylinder, which again is separated by a dark space from the metallic surface. The form which the electric light of the total discharge assumes under the influence of the magnet depends on the quantity of electricity passing at once. The author has specially examined the green light which appears in tubes of common glass at certain degrees of pressure and intensity of the discharge. The luminosity of the sides of the tube is not a phenomenon of fluorescence but of phosphorescence, and can change from green to orange.

§11. The cathodic light which produces this phosphorescence is, as was already assumed by Hittorf, a rectilinear radiation, which extends from the cathode into surrounding space. Still there are essential differences between the diffusion of this remarkable motion and the likewise rectilinear movement of the light, some of which differences are here brought forward.

Hittorf observed that a body placed between the side of the glass and a point-like cathode, throws a shadow in the phosphorescent light of the latter.

Well defined, though not very sharp shadows of small objects, may be obtained not merely from a point-like or linear cathode, but also from extended negative surfaces placed at a small distance from the opaque object.

A surface which merely radiates light, *e.g.*, an ignited body, under similar conditions throws a scarcely visible expanded penumbra.

* The Editor takes the earliest opportunity of bringing before his readers a notice of certain researches due to H. Eugen Goldstein which have hitherto escaped his notice, and which, to a certain extent, anticipate some of the results announced in his papers, "On the Illumination of Lines of Molecular Pressure," read before the Royal Society, December 5, 1878, and "Contributions to Molecular Physics in High Vacuum," read before the Royal Society, April 3, 1879.

The negative light is therefore a rectilinear radiation, which is propagated preferably in a manner almost normal to the producing surface.

§12. Suppose the density of the gas and the intensity of the current in a Geissler's tube so regulated that the side of the tube shows a phosphorescent light round the cathode.

If we introduce along side of and parallel to the cathode a wire not connected with any source of electricity, according to what has preceded, the luminous wire will throw a narrow sharply-defined shadow of the non-luminous wire. As soon, however, as both wires are connected together, and thus converted into cathodes, we perceive on the side of the tube, in the middle of the light-green light, two large, dark, sharply-defined surfaces, which are bisected by a plane passing through both wires. If the wires are straight and not too short their form is oblong, with the longer sides parallel to the wires.

The explanation of these phenomena is:—

The negative electrode is the seat of a repulsive force, which causes every discharge-ray passing near the cathode to turn away.

The figures of the phosphorescent light produced by the cathode-rays are determined by the form of the negative surface. The head of a coin used as a cathode is reproduced in this manner with the accuracy of a portrait in the phosphorescence on the glass.

The author has also studied the stratification of the light. This phenomenon, as regards the positive light, appears in a continuous series of forms, whose ultimate members have little resemblance. The brightness of the successive strata is neither equal nor symmetrical in the direction of the discharge. Each stratum has its maximum brightness near its negative border. Different portions of one and the same stratum differ also in colour.

If we call the "ordinal number" of a stratum the number which shows its successive place, counting from the negative light, the character of every stratum may be pronounced a function of its ordinal number. In all gases the stratification is sharper and more distinct the nearer we approach the negative end of the positive light. This proposition applies not merely to any gas, but to tubes of any, and even compound, form. If we divide a cylinder into a section for the cathode, and into several other sections communicating only by narrow apertures, we find that each stratum of each section has in each other section a stratum corresponding to it in colour, brightness, &c., namely, the stratum having an equal ordinal number. Each single stratum of positive light is an image corresponding to the formerly so-called negative or cathodic light, and the stratified positive light consists of a succession of complexes of negative light.

From the demonstrated identity of positive and negative light, and from the transformation of a negative "bush" into a single positive stratum, we are inversely entitled to regard every complex of negative light essentially as a stratum of positive light.

The boundary of the positive light is independent of the position of the positive pole, and also of the length of the whole discharge.

A displacement of the negative pole in the direction of the discharge effects a displacement of all the positive strata in the same direction. Tubes can be constructed which contain no positive light. With decreasing density all the strata are moved towards the positive pole, while their number decreases.

In a second memoir presented to the Berlin Academy by H. Goldstein, November 23, 1876, the author finds that a portion of the side of the tube, touched internally by a conductor, behaves exactly like a cathode, and emits light possessing all the properties of the cathodic light. The phenomena observed by H. H. Reitlinger and V. Urbanitzky, *i.e.*, a green light opposite a conductor approximated to the tube, are hence merely the excitements which the cathodic rays proceeding from the point of contact must occasion as soon as they, in highly rarefied

gases, extend to the opposite side of the tube. Hittorf has made known in what manner a magnet acts upon the rays of cathodic light. If the results which he has established are applied to the movements of the rays of negative light, there follow, as necessary consequences, all the movements which are observed in the green light under the influence of the magnet. The form of the luminous surface is quite independent of the form of the side touched by a conductor. It can be directly shown with gas-densities somewhat greater than correspond to the production of the green light that the spot touched by the conductor emits a cathodic light. Over the spot in question are seen spheroidal tufts of blue light. The luminous green surface is merely the basis of these blue rays. The author has shown that the positive and negative light can perfectly interpenetrate each other for spaces of any length.

The intensity of the green light emitted from a point of the side of the tube, if the intensity of the exciting radiation is constant, decreases with the continuance of the excitement. This decrease is the greater the stronger the intensity of the exciting rays and the original brightness of the luminous point.

"If between the cathode and the green luminous side of the tube there is introduced a solid body, its shadow is thrown upon the side, since it excludes such rays of the cathode as impinge upon it from reaching the side. If the solid body after some time is removed, the shadow disappears, but an image of the body remains, distinguished from the surrounding luminous surface by its greater brightness, and exactly reproducing the shape of the former shadow."

LOSS OF NITRE IN THE VITRIOL MANUFACTURE.

By JAMES MACTEAR.

THE letters which have been published in recent issues of the CHEMICAL NEWS have thrown considerable light on the above subject, but much still remains uncertain. It seems to me that it is the duty of those who have studied this subject to make public whatever facts they may have ascertained, so that sound conclusions may ultimately be drawn from a careful comparison and consideration of these facts. With this view I beg indulgence and space for some details of the work I myself have done in this direction, and trust it may also induce others to publish the result of their researches.

Shortly speaking, the losses in the case of good plant might be stated as occurring in two ways:—

1. As lost by escaping with the exit gases into the atmosphere.
2. As lost in the acid in the chambers for use (which may or may not have passed through Glovers).

If, however, the amount of nitrous compounds in the escaping gases capable of absorption by caustic soda, and the amount of nitrous compounds in the acid used, be estimated, it will be found that these two sources of loss are very far from accounting for the amount used.

In my researches the estimation of the nitrous compounds in the escaping gases was made by taking average samples over twenty-four hours, the aspiration being at a constant rate (a very necessary precaution); and the determining the nitrogen present in the absorbing solution by means of the zinc and iron distillation with caustic soda process—which I showed in a paper read before the Newcastle-on-Tyne Chemical Society, January 24th, 1878—gives most accurate results when properly carried out.

These daily average samples were checked by an average for the week, and only in very rare cases was there an appreciable difference from the average of the daily tests.

The amount of work involved in these experiments is

very great, but in my opinion it is only possible to arrive at just conclusions by a long series of such tests. The arrangement of these results weekly was as follows:—

Week ending 187

pyrites used.

nitre "

O. V. "

arg. oxygen in exit gases.

" grains NaNO_3 per cubic foot.

vol. per cent NaNO_3 in acid used.

calculated as NaNO_3 on $100\text{H}_2\text{SO}_4$.

NaNO_3 lost.	Tons.	Per cent.	On 100 Sulphur.
In acid used ..	0.228	5.36	0.170
" exit gases ..	0.783	18.36	0.582
As unaccounted for	3.247	76.28	2.418
Total ..	4.258	100.00	3.170

The figures obtained are embodied in the following table, and represent the results of seven series of large chambers, all worked with Gay-Lussac towers, and all but one with Glover towers, the one exception denitrating its acid by hot water in a long tunnel. (See Table on next page.)

It will be seen that the unaccounted for loss has never been under 50 per cent of the nitre used, and in one case has reached 90 per cent, which is not explained by the percentage of nitre used on sulphur, as we have—

Week No. 13, 52.6 per cent of nitre = 2.683 on sulphur.

" No. 23, 90.43 " " = 3.831 "

The total nitre used being—

Week No. 13, 5.10

" No. 23, 4.236.

It would seem that the unaccounted for loss is intimately connected with the working of the chambers, as in most cases it is found that where the percentage of nitre used on sulphur is high, so also is the loss of nitre unaccounted for calculated on sulphur.

As yet no one seems to have hit the blot—Where does the loss take place, and under what conditions? Did we know this we should be able to manufacture our acid at a much reduced cost, were we able to avoid this unaccounted for loss, which is at least 2 to 3 per cent on the sulphur.

The directions in which I have looked for a solution of the problem are—

- A. Decomposition of the nitrous gases in contact with hot kiln gas before entering the Glover towers or chambers.
- B. In their passage through the Glover tower.
- C. In their passage through the series of condensing apparatus generally known as vitriol chambers.

Hitherto my results have been much more of a negative character than otherwise; but a record of failures is of great use, and often leads to a right way out of a difficulty.

I made a number of experiments on a series of vitriol chambers which had no Glover tower, and where during the continuance of the experiments all the nitre was potted in a large pot placed at the end of the row of kilns, all the sulphurous gas passing over it.

The data used in calculating out these experiments in No. 6 series of chambers were—

No. of experiment	hours.
tons pyrites =	sulphur.
lbs. NaNO_3 =	per cent on sulphur.
per cent oxygen in exit gas from Gay-Lussac.	
grains NaNO_3 per cubic foot in inlet gas to Gay-Lussac.	
cubic feet of exit gas calculated from oxygen.	
grains NaNO_3 per cubic foot calculated.	
" " actually found.	
unaccounted for or loss.	
= per cent of NaNO_3 charged =	per cent on sulphur.

LOSSES OF NITRE.

Week	Tons o.v.	Per cent O in exit gas.	Grains NaNO ₃ per cubic ft. Exit Gas.	Vol. P.c. NaNO ₃ .	In Acid used.			In Exit Gases.			Unaccounted for.			Total.		
					Tons.	P. ct.	On 100 S.	Tons.	P. ct.	On 100 S.	Tons.	P. ct.	On 100 S.	Tons.	P. ct.	On 100 S.
1	420	9.0	0.970	0.117	0.613	6.36	0.350	3.448	35.80	1.960	4.572	57.84	3.180	9.633	100	5.500
2	433	10.0	0.940	0.194	1.052	11.30	0.616	3.136	33.70	1.840	5.089	54.70	3.000	9.304	100	5.460
3	443	9.5	0.622	0.102	0.565	5.46	0.327	2.353	22.70	1.362	7.430	71.84	4.311	10.348	100	6.000
4	441	12.0	0.779	0.192	1.034	8.61	0.555	3.865	32.20	2.081	7.113	59.20	3.824	12.012	100	6.460
5	427	10.0	0.915	0.220	1.175	9.24	0.600	3.883	30.54	1.990	7.657	60.22	3.920	12.715	100	6.510
6	440	10.0	1.020	0.156	0.880	7.25	0.450	4.320	35.63	2.210	6.922	57.12	3.560	12.122	100	6.220
7	307	10.0	0.796	0.145	0.552	5.55	0.320	3.025	30.40	1.750	6.373	64.05	3.690	9.950	100	5.760
8	24	10.0	0.452	0.110	0.033	0.55	0.030	0.872	14.40	0.700	5.135	85.05	4.120	6.040	100	4.850
9	440	10.0	0.590	0.190	1.056	12.20	0.700	1.953	22.60	1.300	5.621	65.20	3.750	8.630	100	5.750
10	441	11.0	1.000	0.093	0.529	6.00	0.310	3.509	40.20	2.060	4.696	53.80	2.760	8.734	100	5.130
11	461	10.0	0.730	0.110	0.573	7.00	0.340	2.752	33.30	1.610	4.940	59.70	2.890	8.265	100	4.840
12	463	10.0	0.815	0.075	0.463	6.40	0.300	2.743	37.80	1.770	4.052	55.80	2.620	7.258	100	4.690
13	463	10.0	1.012	0.049	0.287	3.40	0.173	3.710	44.00	2.244	4.420	52.60	2.683	8.417	100	5.100
14	473	11.0	0.777	0.130	0.756	10.00	0.460	2.800	37.00	1.680	4.019	53.00	2.420	7.575	100	4.560
15	470	11.0	0.675	0.061	0.357	5.30	0.216	2.682	39.50	1.616	3.751	55.20	2.260	6.790	100	4.092
16	464	10.4	0.417	0.012	0.069	1.29	0.420	1.543	28.85	0.937	3.736	69.86	2.271	5.348	100	3.250
17	462	11.0	6.497	nil	nil	nil	nil	1.912	34.00	1.185	3.714	66.00	2.301	5.626	100	3.486
18	470	11.5	0.346	nil	nil	nil	nil	1.269	23.03	0.830	4.248	76.97	2.952	5.510	100	3.835
19	437	11.5	0.342	0.058	0.315	6.73	0.258	1.212	25.87	0.877	3.158	67.40	2.255	4.685	100	3.390
20	482	11.5	0.213	0.043	0.270	8.03	0.224	0.659	19.57	0.546	2.437	72.40	2.023	3.366	100	2.793
21	457	11.1	0.239	0.039	0.228	5.36	0.170	0.783	18.36	0.582	3.247	76.28	2.418	4.258	100	3.170
22	437	10.9	0.257	nil	nil	nil	nil	0.915	17.30	0.654	4.375	82.70	3.128	5.290	100	3.782
23	454	10.6	0.113	0.0066	0.036	0.51	0.022	0.384	9.06	0.383	6.568	90.43	3.831	7.290	100	4.236
24	475	8.9	0.432	0.074	0.437	6.82	0.240	1.506	23.52	0.852	4.460	69.66	2.576	6.403	100	3.688
25	403	9.3	0.368	0.032	0.161	2.58	0.093	1.301	20.92	0.753	4.763	76.52	2.759	6.225	100	3.605
26	448	10.4	0.236	nil	nil	nil	nil	0.936	14.80	0.533	5.385	85.20	3.073	6.321	100	3.606
27	463	11.4	0.297	nil	nil	nil	nil	1.418	20.63	0.745	5.455	79.37	2.866	6.873	100	3.611

The following are the results obtained:—

Loss of Nitre in No. 6 Chambers, working large pot (A).

1st expt., 24 hours, 31 per cent of nitre used.
2nd " 60 " 43 " "
3rd " 53 " 68 " "
4th " 43 " 68 " "

Actual average of 7½ days, 60.4 per cent of nitre used.

The average NaNO₃ used during this time was 9 per cent on sulphur and 60.4 per cent of total, or 5.44 per cent on sulphur has disappeared.

During the same time the gases entering Gay-Lussac contained on an average 3.71 grains NaNO₃ per cubic foot, while the amount in the exit gas was 0.30 grain, which shows an absorption of 92 per cent. The gases entering Gay-Lussac were aspirated at a regular continuous rate of 0.5 cubic foot per hour. The pyrites was charged at the rate of 8 cwts. per hour, and the nitre charged varied from 30 to 40 lbs. per hour.

The method of working was now changed, small pots being used for the decomposition of the nitre, and these were placed actually on the burning mass of pyrites in the kilns. The following are the results:—

Loss of Nitre in No. 6 Chambers, working small pots (B).

1st expt., 48 hours, 45.3 per cent of nitre used.
2nd " 48 " 55.4 " " "
3rd " 48 " 42.6 " " "
4th " 24 " 53.7 " " "

Actual average of 7 days 48.6 " " "

Average NaNO₃ used during above time 11.6 on 100 sulphur, 48.6 per cent of which has disappeared, or 5.63 per cent on sulphur.

The gases entering Gay-Lussac contained on an average 4.85 grains NaNO₃, and the amount leaving was 0.058 grain, showing an absorption of 98.8 per cent.

It would appear, then, that as far as these experiments go we have lost in the case where the nitre was decomposed in the large pot in the colder portion of the flue 5.44 parts of nitre for 100 of sulphur, and in the case where the nitre was decomposed in the small pots in the

kilns where the heat was highest 5.63 parts. The difference, 0.23 part, is very small and is not more than the error which might be expected in such experiments, so that in this case I do not think we have proof of a greater loss of nitre by the decomposition of the nitre at a higher temperature, at least not to the extent that might be expected.

On the other hand, there has been required to keep the chambers in as nearly as possible the same condition, in the case of—

Large pot	9.0 per cent of nitre
Small "	11.6 " "

The fact that when using small pots in the part of the flues or in the kilns that more nitre is used is one that is quite beyond doubt and well known to all experienced vitriol manufacturers. It seems to me to depend somewhat on the fact that at the high temperatures the nitrous compounds are driven off much more rapidly, and sent into the chambers, as it were, in "whiffs" or "guats" instead of in an equally distributed manner; it is more difficult to keep the chambers in good condition, and the amount of nitre used is therefore increased.

If we assume that the absorption in the Gay-Lussac be equal in both cases to, say, 95 per cent of the nitrous compounds entering, then the two cases above detailed would show:—

A. 9.0 per cent nitre used—5.44 lost=3.56 entering tower.
B. 11.6 " " "—5.63 " " " =5.97 " "

The loss in exit gases would then be—

A. 0.178 per cent on sulphur.
B. 0.298 " " "

And if we take the exit loss in "A" as 100, the loss in "B" would be 167, or the loss by the method of working as in "B" required 67 per cent more nitre than that in "A," which is sufficiently convincing as to the inadvisability upon this system.

The above experiments, although numerous and interesting, are by no means complete, and more light is greatly wanted.

It is pretty clear that there is a very large loss between the Burners and the Gay-Lussac towers; the experiments in No. 6 series of chambers show that the loss was not very greatly increased by the greater heat at which the nitre was decomposed, and at which consequently the nitrous compounds and sulphurous acid began to react upon each other. We are thus shut up to admit a large loss or reduction of nitrous compounds in the chambers themselves, or no doubt where Glover towers are in use, in the Glovers and chambers combined. So far as my results go there is no proof that the loss is greater where Glover towers are in use; it is, in fact, more the other way, for while the loss where no Glover is used amounts to in—

Case A, 5.44 per cent NaNO_3 on sulphur,

" B, 5.63 " " "

the whole series of experiments extending over more than a year of weekly testings, the loss when Glovers are used is not more than about 3.5 per cent on sulphur. My own opinion is that the greater regularity with which the nitrous compounds are supplied to the chambers has a great deal to do with the question of loss; but this is a point not yet worked out, so far as I am aware, by any one.

The varying statements that one hears made are so contradictory that it were well that those who have trustworthy experiments should publish them, so that by a careful digest and comparison we might come to some valid conclusion. As for myself, I may say that at present I am halting between two opinions, but I trust that experiments now in progress may enable me to decide either for one or the other.

Mr. Davis, in his letter in the *CHEMICAL NEWS*, vol. xxxix., p. 216, refers to the escape of nitric oxide and invites chemists to state their views, after which he will give the experiments which he has made on the subject. As a chemist who has failed completely in obtaining reliable means of determining the amount (if any) of nitric oxide in exit gases, permit me to appeal to Mr. Davis as to whether he should not give us these results at once. As one of the Alkali Act Inspectors, Mr. Davis ought I think to give us any information which may enable us to keep down the escape of noxious vapours, to say nothing of the economy of nitre.

I notice from the correspondence generally that the amount of nitrous compounds present in the actual acid run off for use is not well known. It varies a good deal, but the tests in the table show what it has been in my own case, and the following gives details of a number of tests of the works of my firm at Newcastle:—

Week No.	Vol.	p.c. NaNO_3	in acid	Per cent of Nitre used.
1	0.250			23.0
	0.270	"	"	28.2
	0.205	"	"	23.5
	0.252	"	"	27.4
	0.190	"	"	22.5
	0.108	"	"	11.8
	0.167	"	"	21.5
	0.104	"	"	11.1
	0.122	"	"	13.8
	0.207	"	"	24.0
	0.168	"	"	22.4

St. Rollox, May 20, 1879.

Experiments on Alpine Dairy Farming.—Dr. W. Engling and Dr. v. Klenze.—The vaunted alpine milk differs from other good milk merely by a high percentage of milk-sugar and by the aroma derived from certain odoriferous plants. The influence of manured pastures as compared with unmanured has not been satisfactorily determined, and the authors purpose renewing their experiments. Butter seems to remain free from rancidity the longer the freer the air from ozone.—*Biedermann's Central-blatt.*

ON THE INFLUENCE OF VARIATIONS OF TEMPERATURE ON THE DEVIATION OF POLARISED LIGHT BY SOLUTIONS OF INVERTED SUGAR.*

By P. CASAMAJOR.

(Concluded from p. 214.)

THE announcement that it was at 92° that d became equal to 0, led me to think, that, possibly, the law that I had deduced from Clerget's table was not correct. It is true that the temperatures given by Clerget only extend from 10° to 35°C. , but, between these limits the law which governs the deviations of inverted sugar as affected by temperature, as deduced from his table, is represented by a right line. This I considered as an important fact, because I do not remember a single instance in which a law is expressed by a rigorous right line, between limits that are sufficiently distant, in which this right line is not continued throughout. Having, then, very serious doubts about the accuracy of Clerget's table, I started to discover what the true law was.

For this purpose I prepared pure sugar, by taking the best cut loaf sugar and soaking it in 95 per cent. alcohol for several hours, taking the sugar out, letting the pieces drain, and drying them by hot air.

This sugar, tested before inversion, gave exactly 100 by the saccharometer. The solution, after the direct test, was inverted and tried again in the saccharometer at several degrees of temperature. After making a great many tests at various temperatures, I was forced to the conclusion that not only is Clerget's table correct for temperatures between 10° and 35°C. , but that the law represented by—

$$d = -44 + \frac{t}{2}$$

held good even beyond 88°C. , at which temperature the deviation is equal to 0.

To enable an observer to seize at a glance the results I have obtained, I took a large sheet of cross-section paper divided very accurately into inches and tenths of an inch. On the line of the abscissas I took for every degree Centigrade a space equal to two-tenths of an inch, and, on the line of the ordinates, I took for every division on the negative scale of the saccharometer, four-tenths of an inch. I then drew a right line, connecting the point representing -44 of the saccharometer scale with the point representing 88°C. This right line represents the law expressed by the formula—

$$d = -44 + \frac{t}{2}$$

Afterwards I plotted on this sheet 68 observations made at temperatures varying from 14° to 92°C. The result of this operation was that, out of 68 dots, 10 were entirely erratic, 18 were exactly on the line, and the remaining dots were so close that their distance from the line could be explained by an error of $\frac{1}{4}$ of a division in the observation of the saccharometer. In explanation of these results, I may state that the dots which I have called erratic were due to observations made on turbid solutions, or while the temperature was fluctuating. Observations made under such circumstances cannot be expected to be accurate. The other observations were made under conditions infinitely more trying to the eyes than those under which ordinary saccharometric tests are made. When the temperature fluctuates in the least during an observation, the disk of double quartz becomes distinctly elliptical, and, at times, the line of separation between the two quartz plates swerves alternately to the right and to the left.†

* Read before the American Chemical Society, Feb. 6th, 1879.

† The sheet on which these dots are plotted was shown to the American Chemical Society. I have not thought it necessary to have it engraved, as the explanation given in the text makes the subject sufficiently clear.

To obtain the different temperatures required for these experiments, I had to alter a saccharometer so as to interpose a water-bath between the two optical portions in a manner similar to that adopted by Dr. Ricketts. At the bottom of this water-bath is an opening, communicating with the interior of a closed tube, three inches long, projecting at a right angle. To the closed end of this tube a Bunsen burner was applied when the water-bath was to be heated. In this water-bath I placed the tube containing the solution of inverted sugar. This tube is made of thin brass, and is closed at each end by a glass plate, held by a screw cap in the ordinary way. In the middle of this tube a portion was cut off, two inches long, and as wide as the diameter of the tube. Over the opening thus formed was soldered a projection, as shown in figures 1 and 2 at A. At first I used a tube with a cylindrical projection, of the same diameter as the tube itself, for the introduction of a thermometer to ascertain the temperature of the liquid in the tube, in the manner adopted for Clerget's thick glass tube. I think that the plan represented in fig. 1 and fig. 2 is preferable, as it allows the operator to agitate the solution by moving the thermometer backwards and forwards in the tube. A great advantage obtained by having a projection of this shape is that the

the tube was different from that of the water bath, and that the difference between the two temperatures must depend on the rate of heating the water-bath, *i.e.*, on the size of the flame.

I have mentioned that a thermometer with a long bulb may be the cause of serious errors; so may a thermometer whose movements are too sluggish. In the thermometer that I have used, the bulb was very short; the mercury column was very fine, so as to respond very quickly to slight variations of temperature. These thermometers have 40° C. on a scale six inches long. One goes from 0, to 40°, the next from 30° to 70°, and the third from 60° to 100°. Particular attention was paid in these experiments to keeping the water-bath in continued agitation immediately before taking an optical observation, and while the observation was being made. The thermometer in the brass tube was also kept moving backwards and forwards for some time before looking through the tube. Unless these things are done, there is no certainty that the temperature observed is that of the liquid in the tube.

If, as I am convinced in this case, from the numerous experiments I have made, observations on solutions of inverted sugar, taken at any temperature, are as reliable as those made at any other temperature, there can be no

FIG. 1

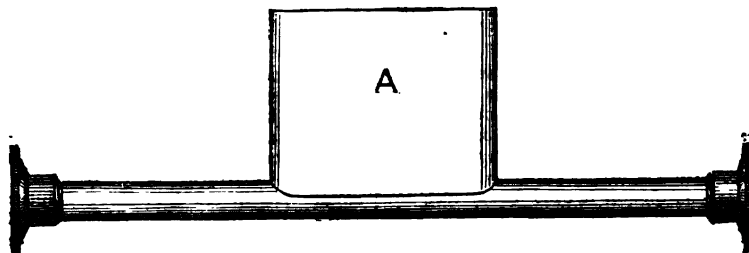


FIG. 2



liquid in the projection does not rise or fall perceptibly when the thermometer is taken in or out. If the vertical projection has a small section, whenever a thermometer is placed in it the liquid rises in this projection, and a portion may be high enough to stand above the level of the water bath. Under these circumstances, this portion of the liquid gets cooler than the portion in the horizontal tube, and, if the bulb of the thermometer is longer than the diameter of the horizontal tube, the temperature indicated will be too low. Too much care cannot be taken in observing these minutiae, as otherwise accurate results cannot be obtained. Even with the greatest care, it is impossible to avoid errors in the observation of solutions whose temperature does not remain constant. In this connection I may be allowed to express the opinion that it was due to the want of a tube like the one represented in fig. 1 and fig. 2 that Dr. Ricketts was led to take 92° as the temperature at which d becomes equal to 0. I have repeatedly found at 92° that the deviation is on the positive side of the scale, ranging from 1.4 to 2, according as the observation was more or less accurately made. In his experiments, Dr. Ricketts did not take the temperature of the liquid in his tube, but that of his water-bath. As his tube was made of glass, about one-eighth of an inch thick, we must suppose that temperature of the liquid in

utility in heating the solution by the interposition of a water-bath. Taking observations at the temperature at which the deviation is equal to 0, merely saves the trouble of dividing—

$$D - d \text{ by } 144 - \frac{t}{2},$$

and it is not certainly worth while to go out of our way for this purpose. The most convenient plan for making observations on solutions of inverted sugar is to follow the directions of Clerget, which are the following:—

A sugar solution is placed in the ordinary way in the saccharometer, and the saccharometric test, D , is noted down. To a portion of this solution, say 50 c.c., are added 5 c.c. of concentrated hydrochloric acid. These are mixed by shaking up the graduated flask, and the flask is placed in a water-bath and heated to 68° C., taking about 10 minutes in raising the temperature. This solution is afterwards immediately cooled to a temperature between 10° and 35° C. We then placed it in a thick glass tube, provided with a vertical tubulure for the insertion of a thermometer to show the temperature of the liquid at the moment of observation. This tube is made 22 centimetres long, instead of 20 centimetres, so as to compensate for the addition of one-tenth of hydrochloric acid. If th

temperature, at the time of observation, is t , and the deviation to the negative side is d , we take the algebraic difference of the two readings, $D-d$, which is the arithmetical sum. To find the correct quantity of cane sugar C , from these observations, we must remember that for 100 per cent. of sugar,—

$$D-d = 144 - \frac{t}{2},$$

therefore,—

$$\frac{C}{D-d} = \frac{100}{144 - \frac{t}{2}} \text{ and } C = \frac{(D-d) \times 100}{144 - \frac{t}{2}}.$$

If we have Clerget's table, we may, instead of making this calculation, follow the practical directions at the head of the table.

Instead of inverting by heating to 68° C., and then cooling down rapidly, it is advised by some authors that the solution be maintained at 70° for at least 15 minutes. I can say, after doing both things repeatedly, that, in the second case, the solution is no better inverted than in the first. Some persons think it necessary to heat at 70° for an hour. This I did not try, as I had found no advantage in heating 20 minutes longer than Clerget directs.

For solutions that are to be heated above 68° for experimental purposes, it is necessary to neutralise the hydrochloric acid by a base, as otherwise the solution becomes very red. For this purpose the preference should be given to carbonate of soda, which gives the best results. I put enough of carbonate of soda to make the solution slightly alkaline, and afterwards make it acid with a slight excess of acetic acid. In this condition the solution may very conveniently be placed in a brass tube. For some reason which I have not been able to discover, when the hydrochloric used in inversion is neutralised by magnesia, the indications of the saccharometer are always too low.

When Venteke's saccharometer is used for testing inverted sugar we are obliged to operate on solutions of greater dilution than the normal solution containing 26.048 grs. of sugar for 100 c.c., because the negative side of the scale is very limited, reaching only to -16 on some instruments. A solution holding 26.048 grs. of sugar in 100 c.c. is tested directly. Then 50 c.c. of this are transferred to the beaker; the 50 c.c. flask is washed out, and the wash water is added to contents of beaker, and also 10 c.c. of hydrochloric acid. After the solution has been inverted and saturated with carbonate of soda, if necessary, the whole is put in a 100 c.c. flask, and enough water is added so that the whole solution at 15° C. shall occupy 100 c.c. The result of the test, after inversion, has then to be multiplied by 2.

I will conclude by giving from my books a series, 28 consecutive tests of raw and refined sugars. In the second column I give the direct test, before inversion; in the third column, the correct test, as afforded by inversion. In a fourth column I have given the copper test for glucose, when I happened to have it. These glucose tests show that the substances which reduce the alkaline tartrate of copper are, for the most part, without action on polarised light.

* Since this paper was read before the American Chemical Society, Dr. Behr has kindly called my attention to a paper of Dr. Tuchschnid in Scheibler's *Zeitschrift* for 1870, p. 649. After a series of elaborate experiments, Dr. Tuchschnid concluded that Clerget's table was very reliable. He investigated the law of this table between 4° and 41.8° C. Instead of the formula given above, he found—

$$R = \frac{100 S}{144.16035 - 0.50578 T}; \quad R \text{ corresponds to } C, S \text{ to } D-d.$$

This formula leads to 87.3° C. as the temperature at which the deviation becomes 0. The law of $d = -44.16035 + 0.50578 T$ is also represented by a right line. As to the results given by this formula, as compared with the one I have given, they may be judged by the following:—

Degrees C.	Tuchschnid.	Clerget.
10°	139.1	139
25°	131.747	131.5
35°	125.458	126.5
4°	123.875	124

Designation of Sugar.	Saccharometric Test (before Inversion).	Corrected by Inversion.	Glucose, Copper Test.
Raw	86½	87	
"	85	86	
Refined—B	91½	92	
" —C	83	83	
Raw	77	80	
"	90½	90½	
"	91	90½	3.57
"	80	84	11.5
"	84	86½	
"	84	89	
"	75½	79	
"	79	81	
"	90	90	4.30
"	95	94½	
Refined	78	78	8.2
"	81	82	7
Raw	92	92	2.5
"	91	91	2.5
"	87	88	5
"	90½	91	2.5
"	88	87	
Manilla	82	81	7.25
"	84	82	7.92
"	83½	81	7.5
Melado	73½	74	7.5
Domestic	82½	81½	
"	85	86	
"	78½	76½	

In the case in which the test before inversion is lower than when corrected after inversion, the presence of an excess of lævo-rotatory substance is indicated. When, on the contrary, the test, before inversion, is higher than after inversion, as in the case of the Manilla sugar, an excess of dextro-rotatory substance is indicated. If these sugars are inverted, and the solutions of inverted sugar are tested in the saccharometer at 85° C., they would show deviations on the positive side of the scale. This, however, does not enable us to decide to what particular dextro-rotatory substance the deviation is due.

COMPOSITION OF A BOILER INCRUSTATION

By ALFRED SMETHAM, F.C.S., A.I.C.

ON analysing a boiler incrustation which was lately submitted to me for examination I found it, after drying at 100° C., to have the following somewhat unusual composition:—

Oxide of iron..	24.72
Oxide of lead ..	8.41
Oxide of zinc..	44.39
Lime ..	0.99
Magnesia ..	0.77
Sulphuric acid ..	1.22
Carbonic acid ..	3.34
Insoluble matter in aqua regia..	5.60
Water of combination, organic matter, and undetermined ..	10.56

100.00

Before committing myself to an opinion as to its cause, I wrote to my clients asking them for particulars of construction of the boiler, and learnt in answer that it was a "hot-water circulating boiler for domestic use," and, further, that the boiler and cylinder were constructed of "galvanised" iron, and communicated the one with the other by means of leaden pipes.

The explanation which I gave of the cause was briefly this:—The town water, with which the boiler was fed, and which is soft to commence with, would, on boiling, be rendered still softer, and by constantly circulating

through the leaden pipes would dissolve a small quantity of the lead. This, passing either into the cylinder or boiler, would be reduced to the metallic state by the zinc with which the iron is "galvanised," and a voltaic action would be induced, the zinc acting as the positive and the lead as the negative metal. By this means a rapid and increasing action would be set up, which after dissolving the zinc would act upon the iron in precisely the same way as the bottom of iron railings, fixed into stone by means of lead, are corroded at their lower ends.

It is thus easy to see how the deposit, which was of a foxy-brown colour and friable, would soon be formed, and the small amount of lime and magnesia salts which are mixed with it prove that the action must have been rapid.

If, as seems probable, there was any metallic lead in the deposit when first removed, this had become oxidised by exposure to the air, to which it was exposed for some time previous to analysis. From a rough sketch of the boiler and fittings which I received, it seems probable that the lead pipes were in metallic communication with the galvanised iron, and if so a much more general action, as well as the local one before indicated, would take place, and destruction would occur at a proportionately greater rate.

Analytical Laboratory,
12, Brunswick Street, Liverpool.

CORRESPONDENCE.

ENAMELLED IRON COOKING-VESSELS.

To the Editor of the Chemical News.

SIR,—I had an opportunity some weeks ago to examine several enamelled iron cooking-pans which were submitted to my approval. By heating the pans two hours on a water-bath, with diluted (7 to 8 per cent) acetic acid and some common salt, and by analysing the solution thus obtained, I found a considerable quantity of oxide of zinc; and as the combinations of zinc are considered to be very injurious to health, and sauces, &c., very often contain mixtures of salt and vinegar, I think it useful to call the public attention to this examination. A family here of seven persons were poisoned by using these pans. The pans were of German and Belgian origin.—I am, &c.,

P. F. VAN HAMEL-ROOS.

Analytical Laboratory, City of Amsterdam,
May 19, 1879.

THE VITRIOL MANUFACTURE.

To the Editor of the Chemical News.

SIR,—Permit me a few lines to point out an error which has crept into Mr. Davis's calculations (CHEMICAL NEWS, vol. xxxix., p. 216).

The error consists in Mr. Davis applying my 20 per cent allowance for chambers working with Gay-Lussac to a system having no such apparatus. When I assumed the loss at 20 per cent I did so fully alive to the fact that the introduction of the Gay-Lussac causes a difference not in the absolute quantity of nitre destroyed per 100 of sulphur, but in the relative quantity, as expressed on the nitre used. In works without Gay-Lussac tower the working nitre and the nitre used are the same quantity, and the loss expressed as percentage on either is identically the same. But in works with Gay-Lussac the working nitre is double and more of the nitre used; hence the same absolute loss will yield in this case two different ratios, depending upon the unit chosen being either the working nitre or the nitre used,—i.e., added to cover losses. When I used the latter quantity as unit for comparison, and admitted 20 per cent of loss in chambers working

with Gay-Lussac, it was equivalent to an admission of a loss of from 5 to 10 per cent in chambers not connected with Gay-Lussac columns.

Thus Mr. Davis's figures would, when in strict accordance with my statement, be as follows:—

	Cwts.
Nitre used	16.0
20 per cent decomposed in chambers, or	
5.3 per cent on the 60 cwts.	3.2
10 per cent loss with exit gas	1.6
10 per cent loss with vitriol	1.6
5 per cent leakage	0.8
Total loss accounted for	7.2
„ not accounted for	8.8

Mr. Davis's own calculation would be more in accordance with Dr. Lunge's statement, since he puts the loss in systems connected with Gay-Lussac towers at 12 cwts. out of 16 cwts. used = 75 per cent of the nitre used.—I am, &c.,

FERDINAND HURTER.

Laboratory, Gaskell, Deacon, and Co.
Widnes, May, 1879.

THE VITRIOL MANUFACTURE.

To the Editor of the Chemical News.

SIR,—Against my wish I must again trouble you, but I hope for the last time on the present question, to afford me some space for replying to Dr. Hurter's and Mr. Davis's letters (CHEMICAL NEWS, vol. xxxix., pp. 215 and 216). So far as the question is concerned in which I took up my pen before, a reply would not be called for; for Dr. Hurter adduces no proof of an appreciable loss of nitre in the Glover tower, merely stating that "he has many reasons for thinking so." I am content to leave that question where it stands, and I refer again to my former letter and to our lengthy discussion in the pages of *Dingler's Journal*. I do not think that there is any inconsistency between my opinion as stated before and that expressed in my treatise, viz., "that certainly the Glover tower is in this respect no worse than any other denitrator," for hitherto, as far as I know, nobody has contended that in the steam columns, &c., any loss of nitrogen takes place by reduction to N_2O or N . Since then, in a paper about to be published, I have proved that denitration by hot water or steam is insufficient when the nitrous vitriol, by faulty work, contains nitric acid.

As I do not wish to extend this letter to an undue length I refrain from going into some fresh matters raised in Dr. Hurter's letter, which are only slightly connected with this issue, much as I could say about them. But I cannot allow myself to be saddled, both by Dr. Hurter and Mr. Davis, with a statement which I entirely repudiate, viz., that the quantity of nitre lost in the chambers, or "in a manner not yet clearly understood" (as Dr. Hurter puts it), is 75 per cent of the total consumption, *whatever that may amount to*. In truth I am not guilty of such an assertion, which, as exemplified by Mr. Davis, involves a downright absurdity. At some factories only 2½ parts, at others—not fitted with recovery apparatus—up to 13 or 14 parts of nitre are used for 100 sulphur; and I am made to say that in either case the loss in the chambers is three-fourths of the total loss! In my letter I had, for the sake of argument, adopted Dr. Hurter's calculation, starting from a loss of 4 parts of nitre on 100 sulphur; and I had proved that, upon his own showing, coupled with evidence referring to the only point in which the two cases differ, the "chemical loss" at works denitrating by steam should be 75 per cent of the above, i.e., 3 parts of nitre to 100 sulphur. Mr. Davis applies that 75 per cent to works where the nitre was not recovered, and where 3 parts were consumed for 100 vitriol, or, say, 9 parts for 100 sulphur, so that my alleged 75 per cent would come to 6.75 parts instead of *three parts*. Dr. Hurter puts the

chemical loss of nitre at works having no recovery apparatus, which notoriously consume 10 to 12 parts of nitre to 100 sulphur, at 20 per cent,—i.e., 2 to 2.4 parts, which is not so very far from my 3 parts. Dr. Affleck, by increasing the absorbing space, has brought down the loss of nitre from 1.45 to 1.05 upon 100 pyrites, or, say, from 3.2 to 2.5 upon 100 sulphur, and he expects ultimately to come down to 0.75 nitre on 100 pyrites,—i.e., 1.67 on 100 sulphur. Even then there might be some loss of nitre in the exit gas as NO. But if we confine ourselves to the consumption of 2.5 parts of nitre, realised by Dr. Affleck as well as by some continental manufacturers (the instance quoted by Mr. Davis, 16 cwts. nitre to 100 tons of vitriol, comes to even less than that), and if we subtract from it the 0.6 part allowed by Dr. Hurter (as 15 per cent of 4 parts) for "mechanical losses" in other ways, the maximum of possible "chemical loss" is reduced to 1.9 parts of nitre to 100 sulphur. But that is only a maximum figure; the real figure, if the exit gas was treated more thoroughly, according to Dr. Affleck might be 1.67, less 0.6 for other mechanical losses, or merely 1 part of nitre to 100 sulphur—not 3 parts, as I had temporarily admitted entirely for argument's sake. In short, it is quite impossible to make even an approximately definite statement of the "chemical loss," and just for that reason I refrained from doing so in my treatise.

I hope that enough mere arguments have been exchanged now. Let us rather try to approach the solution of the difficulty in an experimental way. This I, for my part, have commenced to do in the investigation which I mentioned as being about to appear, and I am continuing my labours in this field.—I am, &c.,

GEORGE LUNGE.

Technical Laboratory of the Federal
Polytechnic Schools, Zürich.

THE CHEMICAL REACTIONS OF THE HOLLWAY PROCESS.

To the Editor of the Chemical News.

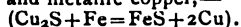
SIR,—The reactions which take place in the Hollway process of rapid oxidation must appear to chemists who have had the privilege of studying the matter to be very interesting (some of those reactions being entirely new), and to be well worth studying.

It is well known, of course, that if iron pyrites is heated in a neutral atmosphere, the result is ferrous sulphide and free sulphur (or something approaching it); but it is not so well known that when ferrous sulphide is heated at a higher temperature, it splits up into a lower sulphide and free sulphur ($2\text{FeS} = \text{Fe}_2\text{S}_3 + \text{S}$). At all events, on blowing through the molten sulphides in a Bessemer converter, sulphur was seen burning abundantly at the mouth, and condensed in the tubes for collecting the gases.

Moreover, as I pointed out at the adjourned meeting of the Society of Arts, the regulus of at least two of the experiments contain sulphur in the proportion to form Cu_2S and Fe_2S_3 .

I.		II.	
Iron ..	57.10	Iron ..	56.05
Copper ..	15.85	Copper ..	16.59
Sulphur ..	21.96	Sulphur ..	23.47

I believe it to be the accepted explanation of the production of moss copper in blue and white metal (at least it is the one given by Percy as the most reasonable) that the subsulphide of iron might be formed, which in presence of subsulphide of copper would form ferrous sulphide and metallic copper ($\text{Fe}_2\text{S}_3 + \text{Cu}_2\text{S} = 2(\text{FeS}) + 2\text{Cu}$); and in connection with this Mr. Schweder finds that by melting subsulphide of copper and metallic iron together the result is ferrous sulphide and metallic copper,—



He also failed to produce subsulphide of iron, and declares that it does not exist.

Of course these results of Mr. Schweder's confirm, to a large extent, what was known or believed by metallur-

gists, but in the Hollway process we find both the subsulphides of iron and copper existing together. It contains no metallic copper, and I have failed to find metallic iron in a similar regulus produced by the same process. Moreover, I consider it extremely improbable, in taking into consideration Mr. Schweder's experiments, that there should be metallic iron and no metallic copper, or so long as subsulphide of copper is present.

It is evident, of course, that the high temperature of the Hollway experiments might be the means of producing the subsulphide of iron, but one would naturally expect that, in cooling, metallic copper and ferrous sulphide would be formed.—I am, &c.,

WM. GALBRAITH.

Sheffield, May 19, 1879.

EXPLOSIONS IN FLOUR-MILLS.

To the Editor of the Chemical News.

SIR,—I noticed with interest the communication of Mr. Watson Smith in the CHEMICAL NEWS (vol. xxxix., p. 222), on the subject of explosions in flour-mills, and as few chemists have had practical experience (as I have had during the last fourteen years) in flour-mills, my observations might not be without interest. I have no doubt from all I have seen that Mr. Smith's explanation of the cause of the explosions is the true one, but I was not aware that he was the first to give it, being under the impression that Dr. Stevenson Macadam had made a very complete investigation immediately after the Glasgow explosion, and arrived at the same conclusions as those now given by Mr. Smith. The remedies Mr. Smith suggests are very difficult to carry into practical operation, as millers require a number of small lights disposed over the mill, and at present the electric light does not conveniently give these. The best plan is to dispense with the stove-rooms altogether, and to filter the air through flannel cloth as it leaves the stone case. This can be done to great advantage by the use of Messrs. Seck's automatic apparatus, which has proved here a perfect success, as not a particle of flour leaves the stone case as dust, mixed with the exhaust air, and the exhaust is greater than without its use could be allowed. There is another source of danger besides that from the stone exhaust, and which it is supposed caused the immense explosion in four mills at St. Louis, U.S.; that is, the exhaust from the middle purifiers, in which the dust carried forward by the air is much drier than that from the stones, and therefore much more easily ignited. In this case also I recommend the filtration of the air immediately it leaves the purifiers. By these means the chance of an explosion is very much less, and should it occur will be much less powerful, as the volume of dust-laden air will be but a small fraction of what it would otherwise be without the filtration process.—I am, &c.,

ARTHUR McDUGALL, B.Sc.

City Corn Mills Manchester, May 26, 1879.

PS.—Mr. Smith suggests the use of an electric lamp in the stove-room. No light is required in that room, with such, if of sufficient volume, need only be entered once per week to be cleared out, and that can be done during the day.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 16, April 21, 1879.

Complementary Researches on the Products of the Distillation of the Alcohols.—I. Pierre and E. Puchot.—The title of this paper is misleading; the

authors treat of the distillation of various commercial forms of ethylic alcohol, and not of the alcohols as a generic group of compounds.

Artificial Preparation of Manganic Peroxide.—A. Gorgeu.—The author produces this compound by heating nitrate of manganese in a glass phial, placed in an oil or paraffin bath, to 155° to 162°. He supposes that in the natural formation of polianite and pyrolute iron suspended in the very fluid mass of nitrate of manganese has been first drawn off, the decomposition of the manganous nitrate taking place subsequently.

On the Tri-tungstates.—J. Lefort.—The author treats of the salts which result from the decomposition of the best known earthy and metallic acetates by sodic tri-tungstate. Equivalent proportions of the two salts are dissolved in a minimum of water. The tri-tungstates are generally instable, and if dried in the water-bath and treated with water they are more or less completely resolved into insoluble bitungstates and soluble quad-tungstates.

Letter to M. Dumas Concerning the Apparatus of Lavoisier.—P. Truchot.—Lavoisier's chemical laboratory and physical cabinet have been reverently preserved by his family, and are now in the possession of M. E. de Chazelles, at Canière, near Aigueperse, Puy de Dome, where the author has seen them. The smallest of the three balances is sensitive to $\frac{1}{10}$ grain. The weights belonging to these balances are wanting, but the kilogramme and its subdivisions as established by Fortin are present, recalling the fact that Lavoisier made all the determinations needful for fixing the weight of the kilogramme. There is a small model of an apparatus for the distillation of sea-water. There are a considerable number of precious stones, some of which have undergone the action of fire. Lavoisier is known to have made an experimental comparison of the heat produced by converging lenses with that of the blowpipe fed with oxygen.

Chemical Function of Anhydrous Acetic Acid.—M. Loir.—Anhydrous acetic acid yields alcohol when treated with a hydrogenising agent. When combined with sodic bisulphite it produces a crystalline compound. With ammonia it forms a crystalline compound insoluble in ether. Finally, acetic anhydride is greedy of oxygen.

On Nitroso-guanidin.—L. Jousselin.—The author prepares this compound by stirring up dry nitrate of guanidin in fuming nitric acid and passing a current of nitrous acid through the mixture. He examines its behaviour with acids, alkalies, and reducing agents.

Value of Certain Chemical Agents Employed in Printing Aniline-blacks.—G. Witz.—The vanadic compounds have been employed for several years in printing textiles. On adding them in extremely minute quantities to thickened mixtures of aniline salts and of chlorates the generation of aniline-black is produced with a rapidity proportional to the doses of vanadium employed. In a letter, published in the *Comptes Rendus* for the session November 25, 1878, M. S. Grawitz has held up chromic compounds as possessing a greater energy than those of vanadium. He founds this opinion upon an experiment in which $\frac{1}{4}$ milligram. bichromate of potassa to a litre of a mixed solution of aniline hypochlorate and of chlorates is said to have produced a colour intense enough to form a kind of ink upon paper. M. Witz has shown that a result such as described, and with such a proportion of chromate was inaccurate (*Comptes Rendus*, December 30, lxxvii.), and subsequently, M. Grawitz, in a further communication, by his silence on this point, seems not to contest the refutation of his original experiment. The modifications introduced by M. Grawitz at the same time, both in the proportions announced and in the nature of the chromates which he now selects in a neutral state (session of February 24, 1879), have given no better results in the check experiments undertaken. It cannot be otherwise, since chrome in its various conditions—and especially

as the chromates of potash, soda, or ammonia—is essentially inert as regards the mixtures of chlorates at the temperature at which the operation is conducted. Here M. Witz considers is the capital error of his opponent, whilst a trace of vanadium acts upon the chlorates in an indefinite manner. The chromates, when in contact with salts of aniline and in a suitable state of concentration, are reduced as soon as the reaction commences, and in proportion as they act they pass into that state of chromic sesquioxide, which remains insoluble and inactive in spite of the presence of the chromates. It is therefore necessary to use quantities of the chromates relatively considerable, because their action takes place only in equivalent quantities. Thickened mixtures containing chromates coagulate rapidly, and to obviate in part this inconvenience M. Grawitz adds ammonia, a device introduced by M. Witz to get rid of any excess of free acid retained among the crystals of aniline hydrochlorate, and which has been well-known and generally used for three years. With this addition, though the development of the colour is slightly retarded, the doctors are completely preserved from corrosion and the design remains clear. When, however, ammonia is used along with the chromates the black is developed suddenly but too much on the surface of the cloth; a certain part of the thickening remains coagulated upon the fibre, and the shade produced has not the intense clear black tone produced with copper or vanadium; it is brownish, imperfect, and much more readily attacked by chlorine. Thus the addition of the chromates to an ordinary colour is distinctly injurious. Even if ammonium vanadate costs 1 franc per gramme it is twelve times less expensive than copper sulphide paste, and certainly much less than the chromates, which act only in equivalent quantities.

Moniteur Scientifique, Quesneville.
May, 1879.

Industrial Society of Mulhouse: Sessions of the Chemical Committee (February 12).—M. Prud'homme described a new colouring matter recently introduced into commerce by A. Bayer under the name of anthracen violet. The author considers that it has no connection with anthracen, but is a substitution-product of gallein, with which it agrees in its reactions, its spectrum, and in the shades produced with different mordants. March 12. —M. Kallab in a note addressed to the Committee admits that his new process for bleaching animal fibres does not yield a white equal in beauty to that produced by the old method, but thinks that the new white, being more permanent, should be preferred where a greenish shade is not objected to. MM. Schmidt and Baldensperger communicated a description of their new processes for the manufacture of magenta by means of vanadium chloride and of certain nitro derivatives, such as nitro-naphthalin. Both yield magenta of excellent quality. Vanadium chloride is allowed to act at 200° upon a mixture of nitrobenzol and commercial aniline, or of pure aniline and nitro-toluol.

The remaining papers in this issue are already familiar to our readers, one dating back as far as 1865.

Biedermann's Central-blatt.
Heft 2, February, 1879.

Belgian Phosphorites.—Prof. Petermann, Crispo, and Mercier.—The enormous phosphatic deposits of Ciply consist of four beds with the following average proportions of phosphoric acid:—

Tuffeau	0.11 per cent
Pierres dures	5.98 "
Craie grise	11.25 "
Poudingue de la Malogne ..	19.75 "

The last mentioned occurs very irregularly and in thin layers, which in many places are already exhausted. The

available quantity of the Craie grise is calculated at 14½ millions cubic metres. Many attempts have been made to render this mineral soluble by steeping in water containing carbonic acid, in solutions of alkaline and ammoniacal nitrates, sulphates, &c., and in the drainage of dung-hills, but the results have been negative. In case of dung-hill liquor the phosphates already existing in solution have been precipitated. The Craie grise has also been applied to soils in the raw state, but without any appreciable effect upon the crops.

Action of Gypsum upon the Quality and Quantity of Clover Crops.—Prof. A. Pasqualini.—In the experiments described the yield of clover was increased, but its nutritive value underwent no improvement.

Excretion of Phosphoric Acid in the Herbivora.—Dr. J. Bertram.—In the carnivora the phosphoric acid introduced into the system is almost exclusively excreted in the urine, whilst in herbivorous animals, under normal circumstances, it appears in the solid excrements, and is not to be detected in the urine unless the lime present in the food is not sufficient for its neutralisation.

Influence of Food on the Quality and Quantity of the Fat of Milk.—Both the percentage and the absolute quantity of fat in milk may vary considerably, even when the diet is unchanged. An addition of protein to the food increases the proportion of fat, but not so much as an addition of oil or stearic acid. The fat of cream melts about 2° lower than the fat remaining in skimmed milk.

Researches on the Ripening of Grapes and other Fruits.—C. Portele.—The ripening of stone fruit after being gathered consists in a decided decrease in acids and in cellulose, whilst the total proportion of sugars is lessened slightly, and in the dextrose is converted into levulose, which is sweeter. There is also under the same circumstances an increase of levulose in grapes, though to a less extent.

Purification of Saccharine Juices with Hydrate of Alumina.—Dr. O. Kohlrausch.—This process, introduced by Prof. Löwig, dispenses with animal charcoal and with the production of treacle. The author finds, however, that a certain quantity of alumina dissolves in the juices, so that the mineral acids present and also the colour are not entirely removed. Hence it is very doubtful whether the filtration over animal charcoal could be omitted. The production of treacle is also not obviated in Löwig's process. The elutriation of the hydrate of alumina requires

such quantities of liquid that the boilers and presses would require to be enlarged, and the consumption of fuel would be increased.

Foreign Colouring Matters in Red Wines.—Prof. J. Nessler.—The author considers that we are not yet able to distinguish the red colouring matters of mallows and bilberries from that of grapes. It is very probable that they do not essentially differ.

On Casein.—Prof. O. Hemmarsten.—The author prepares casein by precipitation with acetic acid, avoiding excess, re-solution of the washed precipitate in a minimum of alkali, so that the liquid retains a faint acid reaction, filtration to remove fat, re-precipitation with acetic acid, and after several repetitions of the above-described process washing with alcohol and ether. Pure casein possesses the characters of an acid; if dissolved in a minimum of caustic alkali it has an acid reaction; it dissolves the carbonates of lime and baryta with liberation of carbonic acid, and even phosphate of lime.

Heft 3, March, 1879.

Temperature of the Soil in a Compact and a Loose Condition.—Prof. E. Wollny.—In summer and in warm weather compact soils are on the average warmer than loose ones, but in winter, and on a fall of temperature in summer, they are colder. In warm weather compact soils are warmer by day and colder by night than loose ones, and are subject to greater fluctuations of temperature.

Composition of the Milk of Cows of Different Breeds.—E. Marchand.—The author finds free lactic acid, a constituent hitherto invariably overlooked, present to the average amount of 2 grms. per litre. The relative proportions of albumen and casein fluctuate much, but their joint weight is little affected. If the cow has eaten cruciferous plants the milk is rendered poor in casein and rich in albumen, thus effecting the production of cheese.

Composition of Clover-hay Rapidly Dried or Exposed to Rain.—Dr. C. Brimmer.—The quantity of protein is reduced 0·7 per cent, but the non-nitrogenous extractive matters are diminished by 7·1 per cent.

Artificial Horse-dung.—Prof. Mayer.—A manure is said to be exported from England to Holland which has the outward appearance of stable manure, but on examination proved to be a mixture of straw and decayed leaves.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

APRIL, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

Appearance in a foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	Sulphuric An- hydride.	Hardness on Clark's Scale.	
	Grains.	Grains.	Grains.	Grains.	Grains.	Grains.	Grains.	Grains.	Grains.	Before Boiling.	After Boiling.
<i>Thames Water Companies.</i>											
Grand Junction	Clear	0·000	0·005	0·135	0·026	19·70	7·700	0·670	1·220	1·430	14·8 3·30
West Middlesex	Clear	0·000	0·003	0·105	0·015	20·90	7·590	0·720	1·880	1·600	14·8 3·30
Southwark and Vauxhall	Clear	0·000	0·006	0·126	0·047	19·80	7·590	0·720	1·880	1·670	14·3 3·30
Chelsea	Clear	0·000	0·008	0·114	0·052	19·70	7·630	0·460	2·000	1·660	13·7 3·30
Lambeth	Clear	0·000	0·008	0·135	0·044	20·70	8·150	0·610	1·880	1·570	14·3 2·80
<i>Other Companies.</i>											
Kent	Clear	0·000	0·003	0·315	0·005	28·10	10·860	0·841	2·830	2·970	17·6 5·10
New River	Clear	0·000	0·003	0·135	0·008	21·50	7·950	0·540	1·880	1·130	14·3 3·30
East London	Clear	0·000	0·006	0·165	0·028	21·40	8·230	0·684	2·120	1·970	14·8 2·80

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYNOTT TIDY, M.B.

NOTES AND QUERIES.

Manufacture of Sulphuric Acid.—Has it ever been suggested that the pot containing nitre or nitrate of soda and sulphuric acid be replaced by one containing sulphur and nitric acid, so that the manufacture of a by-product, and the heat, time, and labour consequent thereon be avoided. Of course this plan would not be so suitable for stone acid manufacturers. The principal objection would seem to be the greater cost of the nitric acid, but when the densities of the former and sulphuric acid are taken into account, the cost is considerably lessened.—**SULPHURIC ACID, Swansea.**

Mineralising Flannel.—Can you or any of your readers inform me of a process by which flannel may be mineralised or preserved from the action of a continuous flow of water at a high pressure in which it is immersed, as I find that after a few months use the flannel loses its nature, the threads being partially decomposed? Would water have the same effect upon silk?—**ANDREW BELL.**

Blue Manifold Paper.—Will you kindly favour by answering, through the columns of your valuable paper, a recipe for making blue manifold paper.—**B. M. P.**

MEETINGS FOR THE WEEK.

- MONDAY, 2nd.**—Royal Institution, 3. "The Intellectual Movement of Germany from the Middle of the Last to the Middle of the Present Century," Prof. Hillebrand.
General Monthly Meeting, 5.
- TUESDAY, 3rd.**—Royal Institution, 3. "Suggestions to Students and Readers of History," Prof. J. R. Seeley.
Zoological, 8.30.
- THURSDAY, 5th.**—Royal Institution, 3. "Suggestions to Students and Readers of History," Prof. J. R. Seeley.
Chemical, 8. "On Eardennine," Dr. Stenhouse and Mr. Groves. "Theory of Fractional Distillation," F. D. Brown. "Action of Organo-zinc Compounds on Quinones," F. R. Japp. "On Chlorotannic Acid," J. W. Mallet, F.R.S. "On Indigo-purpurin and Indirubin," Edward Schunck, F.R.S. "Third Report to the Chemical Society on some Points in Chemical Dynamics," Dr. Wright, Mr. Luff, and Mr. Rennie.
- FRIDAY, 6th.**—Royal Institution, 9. Prof. Dewar.
Geologists' Association, 8.
- SATURDAY, 7th.**—Royal Institution, 3. "On Swift," Prof. H. Morley.

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THE CHEMICAL NEWS.

VOL. XXXIX. No. 1019.

9 JUN 79

NOTE ON THE SPECTRUM OF SODIUM.

BY J. N. LOCKYER, F.R.S.

I HAVE lately been engaged in studying the spectrum of sodium under new experimental conditions. In anticipation of a detailed communication I take leave to state that the vapour given off from the metal after slow distillation in a vacuum for some time shows the red and green lines without any trace whatever of the yellow one. Hydrogen is given off in large quantities, and at times the C line and the red "structure" are seen alone. After this treatment the metal, even when red hot, volatilises with great difficulty.

PRELIMINARY NOTE ON THE ABSORPTION OF SELENIUM BY PLANTS.*

By CHARLES A. CAMERON, M.D.,
Professor of Chemistry and Hygiene, R.C.S.I.

The subject of the possibility of replacing some of the elements found in plants by other elements of the same atomicity has not engaged the attention of British chemists; but on the Continent a few attempts in this direction have been made, generally with but little success. For example, Berner and Lucanus vainly attempted to replace ferric oxide (Fe_2O_3) in plants by the manganese analogue of that compound (Mn_2O_3). Experiments made with the view of substituting sodium for potassium in plants have invariably given negative results. The possibility of completely replacing an element in plants by another was, however, proved by me in a paper read before this Society in 1863. I found that rubidium was capable of taking the place of potassium. It may be that certain bodies, though not capable of completely replacing other substances in vegetables, may be partially substituted therefor. The varying proportions of sodium and potassium found in the ashes of plants would seem to indicate such a partial replacement. As a rule, whenever potassium is sparsely present in the ashes of a plant, sodium abounds therein, and *vice versa*.

The analogy between sulphur and selenium suggests the possibility of the latter wholly or partly replacing the former as a constituent of vegetables. Sulphur exists in plants on two conditions, namely, as a constituent of sulphuric acid, and as an ingredient of albuminous and certain oily bodies. Sulphur is only taken up into the mechanism of plants in the form of sulphates, such as, for example, calcium and sodium sulphates. By the partial deoxidation of these sulphates, sulphur is procured by the plant and employed in the elaboration of its albumen, casein, and other albuminoids. The results of the interesting experiments of Arendt render it probable that sulphur is both oxidised and deoxidised in the plant at different periods of its development. Thus, after blossoming the oat was found to contain more sulphur trioxide than before the ears of the plant had formed; and whilst the sulphur trioxide disappeared altogether from the lower part of the stem, it increased largely in the leaves and in the plant at large.

Now, there are two problems to be solved in relation to the substitution of selenium for sulphur in plants. First, can selenium in the form of selenic acid replace sulphuric

acid? Second, can selenium replace sulphur as a constituent of albuminous bodies? If it were possible through the agency of the organs of plants to effect the replacement of an element not merely in an organic, but in an organised body, the achievement would be one of the greatest importance. Absolutely nothing has been done in the way of effecting the synthesis of new bodies by means of the wonderful combining powers which are exerted in the vegetable mechanism.

The difficulty with which compounds of selenium analogous to the organic bodies containing sulphur are prepared, and the instability of so many of them, are facts which deter one from feeling sanguine as to the possibility of effecting the synthesis of organised selenium bodies by means of plants; still the attempt is worth making. Much more hopeful is the chance of replacing sulphuric acid by selenic acid. I am of this opinion from the results of an experiment which I made several years ago upon a very small scale, and which I never published up to the present, every year intending to repeat it upon a larger scale. The experiment was as follows:—A sod was taken from a field in which a crop of the so-called artificial grasses (which are chiefly leguminous plants, and not grasses at all) was just peeping over ground. The sod was 2 feet in depth, 3 feet in length, and 1 foot wide. It was placed in a box, and one-half of the plants were watered twice a week with a weak solution of potassium selenate (K_2SeO_4). During four weeks the total quantity of potassium selenate applied to the plants amounted to 20 grms., which comprised my whole stock of the article.

Now, this experiment was merely a tentative one. First, to ascertain whether or not selenic acid would act injuriously upon plants. Secondly, to discover whether or not the selenic acid could partly replace the sulphuric acid, or rather could be taken up into and permanently retained by the plant. With respect to the action of the selenate, I could not perceive any difference between the plants to which the salt had been applied and those to which it had not. I came to the conclusion, therefore, that selenic acid applied at least in small quantities did not injure plants.

The next step was to ascertain whether or not the selenic acid had been partially absorbed. The plants were accordingly partially dried, and boiled in strong nitric acid until thoroughly destroyed. The solution was evaporated to dryness, and the residue was treated with dilute hydrochloric acid, which dissolved it nearly completely. The solution was concentrated and mixed with a saturated solution of sulphurous acid, whereupon the liquid assumed at once a deep blood-red colour, from the separation of selenium.

Before the plants were partially dried they were carefully washed, in order to separate any traces of selenate which might have adhered to their surface. The application of the selenate solution was discontinued for a fortnight before the plants were taken up.

I think this experiment proves that selenic acid is not injurious to plants when used in small quantity, and that the acid is taken up and retained by plants, or at least of certain varieties of plants. The experiment, however, did not prove whether or not there was a partial replacement of sulphur trioxide by selenium trioxide or of sulphur by selenium. Having lately become possessed of large quantities of selenium compounds, I proposed to grow plants in soil or water free from sulphur in any form, but supplied with potassium and ammonia selenates. Should the results of this proposed experiment prove interesting I shall do myself the honour of submitting them to the Society.

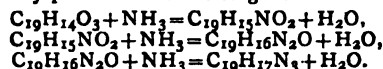
Proportion of Carbonic Acid in the Atmosphere.—J. Reiset.—The air contains in 10,000 parts 2.942 parts of carbonic acid by volume. The most extreme variations have not exceeded 3 parts in 10,000.—*Comptes Rendus*.

* Paper read before the Royal Dublin Society, May 19, 1879.

ON AURIN.* PART II.

By R. S. DALE, B.A., and C. SCHORLEMMER, F.R.S.

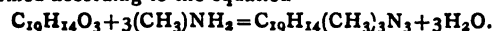
SOME time ago we read a paper before this Society, in which we stated that on heating aurin, $C_{19}H_{14}O_3$, with an excess of aqueous ammonia, it is converted into parosanilin, $C_{19}H_{17}N_3$.† There ought to exist two compounds standing intermediate between aurin and rosanilin, viz., $C_{19}H_{15}NO_2$ and $C_{19}H_{16}N_2O$, as the action of ammonia probably proceeds in three stages:—



We have made a very large number of experiments for the purpose to get hold of one of these intermediate products, and have at last succeeded in obtaining one, probably the first, in beautiful crystals; it dyes on silk and wool a rich red, and as the commercial red aurin, which is obtained by the action of ammonia on common or yellow aurin, has also been called *peonin*, we will retain this name for our compound. It is formed by heating aurin with dilute ammonia to $100^\circ C$. for some weeks. A body dyeing exactly the same shade, and therefore undoubtedly the same compound, is obtained by passing for a few hours ammonia gas into a solution of aurin in boiling amyl alcohol.

As we encountered many difficulties in the preparation of peonin, we thought more definite results might be obtained by using a compound ammonia, and therefore tried the action of methylamin on aurin. Methylamin is now a commercial article, being obtained from beetroot-molasses by M. Vincent's method, which Dr. Roscoe described to this Society some weeks ago; he was kind enough to present it with the methylamin required for our experiments.

On heating the two bodies together for a few hours, the deep red colour of the liquid became much paler, and a brown deposit was formed in the tubes, which on examination was found to be *trimethyl-para-rostanilin*, which is formed according to the equation—



We have dyed with it, as well as with the other bodies mentioned in this paper, silk, wool, and mordanted cotton, and beg to submit these samples to your inspection.

We have already proved that by acting on aurin with aniline the final product consists of triphenyl-parosanilin, and thus shown that the whole host of bodies known as the aniline colours can be obtained from phenol.

ON THE COMPOSITION OF THE ASHES OF WHEAT- BRAN, BURNED UNDER A STEAM-BOILER.

By S. F. PECKHAM.

SOME time ago, at a meeting of the Minnesota Academy of Natural Sciences, a substance was handed me by someone present with the remark that it was the fused ashes of wheat-bran, burned under a steam-boiler. The material was nearly pure white in colour and vesicular in structure, resembling more nearly some specimens of Cooper River phosphate than anything else that I remember. Wishing to learn its composition I submitted it for analysis to Miss Cora J. Brown, then a student in the University Laboratory. Miss Brown had already proved herself a very careful and skilful manipulator, and she conducted the analysis with exquisite neatness and patience, and with the most gratifying results.

The material had the appearance of having been completely fused, and was very free from even minute specks of either charcoal or sand. It had a specific gravity of 2.34, and a hardness of 3.5 to 4.

The analysis gave the following results:—

	Per cent.
Chlorine	0.612
Sulphuric oxide (SO_3)	0.151
Silicic " (SiO_2)	0.627
Phosphoric " (P_2O_5)	48.328
Potassium " (K_2O)	6.659
Sodium " (Na_2O)	6.664
Calcium " (CaO)	10.615
Magnesium " (MgO)	18.980
Ferric " (Fe_2O_3)	2.015
Water	0.438
Sand	3.170
	98.259

Which may be interpreted as indicating the following composition for the ash:—

	Per cent.
Potassium chloride (KCl)	1.2887
" silicate (K_4SiO_4)	2.5936
" phosphate (K_3PO_4)	5.8337
Sodium phosphate (Na_3PO_4)	11.7370
Hydrogen phosphate (H_3PO_4)	9.3721
Calcium sulphate ($CaSO_4$)	1.9567
" phosphate ($Ca_3(PO_4)_2$)	18.2342
Magnesium phosphate ($Mg_3(PO_4)_2$)	41.4600
Ferric phosphate ($Fe_2P_2O_8$)	3.8058
Water, hygroscopic (H_2O)	0.4379
Sand, insoluble residue.. .. .	3.1700
	99.8897

The remarkably large percentage of phosphoric oxide present at once attracted my attention, and at my suggestion the determination was repeated. The results of the four determinations made with uranium acetate were very closely concordant, and I think of unimpeachable accuracy.

No carbonic oxide (CO_2) was present. The very small proportion of chlorine is doubtless due to the high temperature to which the ash had been subjected—a temperature sufficiently high to completely fuse it. The substance may therefore properly be considered an ash residue consisting of its less volatile portions. The presence of free phosphoric acid in such a compound may be questioned; perhaps also the presence of normal magnesium phosphate. It must be remembered, however, that it is scarcely a supposable case that the phosphates in the bran are acid phosphates, producing pyrophosphates when burned; and while it would appear more probable that the albuminoid phosphorus contained in the bran would burn into phosphoric oxide and escape in the smoke, it is not a necessary supposition, especially when the results of analysis show that there is more phosphoric oxide present than is sufficient to form normal phosphates with the available bases. I have therefore assigned the composition given above as the most satisfactory interpretation of the results.

Laboratory of the University of Minnesota,
May 7, 1879.

Heat attending the Formation of Cyanogen.—M. Berthelot.—The mean heat evolved on the combustion of cyanogen, calculated for its equivalent $C_2N=26$, is 132.3 calories. In its formation heat is absorbed to the extent of 38.3 cal. Hence, like acetylen and nitric oxide, substances which play the part of true compound radicals, it is a body formed with disappearance of heat.—*Comptes Rendus*.

* Paper read before the Manchester Literary and Philosophical Society, April 2, 1879.

† Mem. Lit. Phil. Soc. (3), vi., 248.

NOTE ON THE DETECTION OF SOME
RARE METALS IN PYRITES FLUE DUST.*

By DAVID PLAYFAIR,
Assistant in St. Rollox Laboratory, Glasgow.

THE flue dust which was examined was taken from a short flue leading from pyrites kilns in which Spanish cuprous pyrites were burned. As the flue in question was an iron tube not more than 2 or 3 yards long, anything that condensed in it must have done so at a very high temperature; and that probably accounts for the very small quantity of selenium found, as that metalloid is found in appreciable quantities in the sulphuric acid manufactured from these pyrites.

The greater part of the flue dust consisted of compounds of arsenic, antimony, lead, copper, and iron; but it is only the thallium, tellurium, and selenium that this note is concerned with.

Thallium.—The flue dust was treated with sulphuric acid for the extraction of the thallium, which was precipitated as chloride. The chloride was re-dissolved and re-precipitated twice, and then reduced to the metallic state with cyanide of potassium. One sample of flue dust, treated thus, gave about 0.05 per cent of thallium, while in another case the amount obtained did not exceed 0.002 per cent.

Tellurium.—The insoluble matter left after the above treatment with sulphuric acid was boiled in a solution of caustic soda. The solution was filtered off, acidified with hydrochloric acid, boiled, cooled, saturated with sulphurous acid, and boiled again, when the selenium and tellurium were precipitated. This precipitate was collected, dried, and then fused with cyanide of potassium in an atmosphere of coal-gas. The fused mass was allowed to cool, and then dissolved in warm water saturated with coal-gas, and through which a stream of coal-gas was kept continually passing, on a filter. The telluride of potassium thus dissolved passed through the filter exhibiting its characteristic claret colour, which it, however, immediately lost, becoming colourless, and precipitating tellurium unless a stream of coal-gas was kept passing through the filtrate also.

The amount of tellurium found was about 0.002 per cent of the matter undissolved by sulphuric acid in the treatment for thallium. The amount of selenium did not exceed 0.001 per cent.

CHEMICAL JOTTINGS.†

By H. R. PROCTER.

I. *Weselsky's Reaction for Phloroglucin.*

THIS reaction is described in the *Chemical Society's Journal*, 1876, i., 964, and consists in the production of a cinnabar-red precipitate when a trace of phloroglucin is added to a solution of nitrate of aniline or toluoydin containing a little nitrite, and the mixture allowed to stand some hours.

It is known through the researches of Hlasiwetz (*Ann. der Chemie*, cxliii., 290) and others, that many tannins when heated with dilute sulphuric acid are resolved into glucose and peculiar red bodies which we may call phlobaphenes. These again, when fused with potash, are decomposed into protocatechuic acid and another body, which in some cases is a fatty acid, and in others as the species of sugar known as phloroglucin.

To be able to recognise these phloroglucide tannins without the tedious process of separation noted above would be an important aid to the classification and qualitative

identification of these complex bodies; and it occurred to me that possibly they might give Weselsky's reaction in their unaltered condition. To test the matter, I mixed in test-tubes 5 c.c. water, 1 c.c. of a saturated solution of nitrate of aniline (probably containing toluoydin), and 1 c.c. of very dilute solution of potassic nitrite, and added to each tube 1 c.c. of solutions containing as nearly as possible 5 grms. per litre of the following bodies:—(See Table, next page.)

In five minutes most had become yellow, gambier, and oak, and churco barks deeply so, and in an hour oak and churco barks had deposited a deep red precipitate. In twelve hours all had precipitated more or less, but the glucose and sugar yielded only traces. The gambier, which contains catechin, a well known phloroglucide, gave a fine deep red precipitate, and oak and churco barks, which very probably contain phloroglucin, gave precipitates of which the lower part was equally bright.

It seems evident, therefore, that phloroglucides in combination may yield this reaction, but that many other bodies also give reddish-brown precipitates which may easily lead to mistaken conclusions. When more dilute solutions are used the same facts are observed, but the red precipitate is more distinct from the rest. Some precipitate was even obtained on long standing from aniline nitrate and potassic nitrite alone. The precipitates seem remarkably stable, being unaltered by solution of iodide, dilute acids, ammonia, sulphurous acid, or sulphide of sodium.

The yellow supernatant liquid, if acidified and exposed to the air, becomes bright crimson, and sometimes deposits minute crimson double-refracting acicular crystals. It is turned yellow by ammonia and its colour restored by acids.

Watts states that pyrocatechin gives Weselsky's reaction.

The churco bark to which I have alluded is the product of the *Oxalis gigantea* of Chili, and contains a red tannin equal in amount, by Löwenthal's test, to from 25 to 26 per cent. quercitannic acid. The sample in my possession was kindly supplied by Messrs. Machado, of Glasgow.

I have made some preliminary experiments as to the application of Wiesner's test for woody fibre—the purple colouration produced on moistening it with solution of phloroglucin and then with the hydrochloric acid (*Chemical Society's Journal*, 1878, p. 809)—to the converse detection of phloroglucides. Deal shavings moistened with solution of gambier gives it strongly. I hope to pursue the subject further.

II. *Explosive Product of Solution of Phosphorus in Bisulphide of Carbon.*

In packing my chemicals for removal to a new laboratory, I came upon two bottles of this solution, probably five or six years old, which had formed an orange-yellow deposit on the sides. Having recently read of a case, to which I am sorry that I cannot at the moment refer, in which a bottle with a similar deposit had exploded violently on being broken, I determined to try the experiment at once, lest it might try itself under more dangerous conditions. The bottles were thrown against a wall, and one exploded with a large puff of flame and white smoke: the other gave a slight flash, and smoke which smelt rather of phosphoretted hydrogen than of phosphoric acid. It is evident that some new and spontaneously inflammable compound had been formed, since the fresh solution only inflames after some minutes, when the carbon disulphide has had time to evaporate, and fails to do so at all when the temperature is very low. In the present case the thermometer was below freezing-point, and combustion was instantaneous. Phosphoretted hydrogen is out of the question, unless moisture had found its way into the bottle.

Of course I can claim no originality in this observation, but think it well to put it on record as a caution, sir—

* Read before the Chemical Section of the Glasgow Philosophical Society.

† Read before the Newcastle-upon-Tyne Chemical Society, March 27, 1879.

No.		In Five Minutes.	In One Hour.	After standing over night.
1.	Gallic acid	Yellow colouration	Orange, no precipitate	Dark red-brown precipitate
2.	Gallotannic acid	Ditto	Ditto	Ditto
3.	Gambier	Deep yellow	Orange, turbid	Venetian red precipitate (compact)
4.	Oak bark tannin (50 grms. bark per litre)	Ditto	Orange, red precipitate	Ditto, upper part darker
5.	Pyrogallol	Pale yellow	Orange, turbid	Reddish brown flocculent ppt.
6.	Churco tannin (20 grms. bark per litre)	Deep yellow	Dark orange, red precipitate	Red precipitate, flocculent, dark above
7.	Cane-sugar	Colourless	Clear yellow liquid	Trace brown precipitate
8.	Impure glucose	Ditto	Ditto	Ditto

the reaction might easily give rise to a dangerous accident.

III. Determination of the Free Acid in Tan Liquors.

Tan liquors usually contain, in addition to tannins, gallic acid, catechu, and other bodies of very feeble acid properties, a certain portion of acetic and other fatty acids, and frequently free sulphuric and hydrochloric acids, which have been purposely added. For technical purposes it is often desirable to determine these stronger acids, which are capable of dissolving lime and swelling the tissue of hide in presence of tannin, &c., which have the opposite effect. This cannot be done by an alkaline solution and litmus, since the tannins are immediately oxidised in alkaline solution, giving rise to dark coloured products, which obscure the end-reaction.

A simple and sufficiently accurate method for technical purposes is to use lime-water as the standard solution, adding it to the clear filtered liquor in a beaker until permanent turbidity is produced. This point is easily recognised by holding the beaker over a printed page. The end reaction, of course, depends on the fact that calcic tannates are soluble, or more probably cannot be formed in the presence of a free and stronger acid. I have not attempted to determine exactly *what* acids are estimated by this process, the mixture is complex, and the amount of lime which they are capable of dissolving is the essential point in practice.

REPORT ON THE EXPERIMENTS WITH THE ELECTRIC LIGHT ON THE VICTORIA EMBANKMENT.

THE joint official report of Sir J. W. Bazalgette, the engineer, and Mr. T. W. Keates, the consulting chemist of the Metropolitan Board of Works, has recently been made public, and is a model of what such a report should be.

For the first time in the history of the electric light, as applied to street illumination, we have a series of data laid before us about whose accuracy there can be no doubt. Hitherto such reports have emanated from either gas-managers or electricians, whose interests—wrongly we think—have been held to be opposed to each other; hence the singular discrepancies in the data which have been issued up to the present time. It must be perfectly understood that we in no way intend to make the slightest imputation against the honour of the authors of these reports; but when one side tells us that the electric light is cheaper than gas, and the other that gas is cheaper than the electric light, we can only account for the discordance of opinion by supposing that the reporters' judgment must have been unconsciously influenced by their anterior convictions. In the present instance we have the conclusions of two eminent public officials, who have no other interests to serve than those of the community at large, and who are wholly independent of the advocates or opponents of either system of lighting.

We regret that we cannot follow the reporters through all the elaborate series of calculations which they have laid before us; we can only gather a few of the results of

their carefully conducted experiments. One great point which has been decided on the Victoria Embankment is that the steadiness of the light is in a great measure dependent on the regularity with which the motive power is supplied to the electric machine. In the present case this uniformity of power was secured by the use of a specially constructed 20 horse-power engine, provided with an automatic governor of extraordinary sensitiveness, the whole being supplied gratuitously for three months by Messrs. Ransomes, Sims, and Head, of Ipswich.

The experiments upon which Messrs. Bazalgette and Keates's figures are founded began on January 24th, and were continued until February 5th, Sundays excepted, — or, in other words, for twelve nights,—the engine being driven for 5½ hours each night, the whole 26 lamps being lighted. The mean daily cost of working the engine, including wages, was found to be £1 9s. 8½d., or 3¼d. per lamp per hour. The cost of the whole plant was £1280, but a smaller engine would be quite sufficient for the purpose; the capital required, therefore, might be reduced to £990.

"The accurate estimation of the amount of light given off by the electric arc and ignited carbons is," say the Reporters, "a matter of great difficulty, so much so that the best results which have been obtained can only be looked upon as fairly approximating to the truth." This difficulty arises from two causes: first, from the difference in colour between the electric light and any standard of light at our command; and secondly, the fluctuating character of the electric light. The photometrical experiments made on the Embankment were—1st, on the naked light; 2ndly, on a light with an opaline globe; and 3rdly, on a light with a granulated globe. The measurements were made with an ordinary bar photometer, fitted with a movable screen with a star disk.

The standard unit of light which was used was a sperm oil lamp, specially constructed for the purpose by Mr. Sugg, and fitted to the photometer by a peculiar balance, by which the necessity for removing the lamp during the course of any number of experiments is obviated. The unit of light given by this lamp was equal to 16 Parliamentary standard sperm candles. Twenty readings were made on each experiment, both with open and shaded lights,—that is to say, 160 on the naked light, 24 with the opaline globe, and 16 with the granulated globe. The following are the mean results obtained in these experiments:—

Naked light	378.1 sperm candles.
Opaline globe	154.9 " "
Granulated globe ..	265.0 " "

The loss of light produced by the opaline globes is 59 per cent; by the granulated globes, 29.9 per cent.

Taking the above data as approximately correct, the gross output of light from the twenty Jablochhoff candles amounts to 7562 candles, nearly 60 per cent of which is absorbed at once by the opal globes. There are also two other sources of loss: first, from the greater thickness of the opal glass in the lower third of the globes; and secondly, from the fact that the most favourable direction for the projection of the light from the Jablochhoff candle is upwards, there being an actual diminution of light horizontally, and still more so downwards.

On the 14th and 15th of March a very important expe-

riment was made, with a view of ascertaining whether the circuits could be increased in length to a considerable degree without diminishing the apparent value of the lights on such circuits. The Westminster and Waterloo circuits—that is to say, the two at each end of the line—were thrown into one, the total length of wire being 7063 feet, three of the lights being at the Westminster end and the other two at the Waterloo end. The lights burned with apparently equal brilliancy as when short-circuited, and on the following day the length of the line was increased to 8815 feet, or 1.66 mile, but still without any perceptible effect on the light. We believe this is the longest circuit ever attempted to be used with an alternating current machine, and the success of the experiment has a very important bearing on the general question of electric lighting.

We have said before that the cost of a plant necessary for working twenty Jablochkoff lights is £990. The interest on this at 5 per cent would be £49 10s., and taking wear and tear at 10 per cent, £99, we get a total of £148 10s. Taking the whole time during which the lamps are lighted in the year at 3600 hours, say 10 hours a day, the cost of use of plant for each light per hour is 0.49d. To this we must add 3.24d. per light per hour for the cost of producing the light, and 2d. per hour per candle, making a total of 5.73d. per light per hour all the year round. Let us now compare the cost of gas with the latter figure. The light in the opal globe, as given above, is in round numbers equal to 155 candles, and in the granulated globe to 265 candles. To produce a light equal to 125 candles, one of the larger Sugg burners consumes 48 cubic feet of gas per hour, while to produce a 265-candle light we should require 83 cubic feet per hour. The cost of the former quantity of gas may be taken at 2d., of the latter at 3.5d., or, to put the matter more plainly,—

The cost of the electric light is ..	5.73 per hour.
Gas equivalent to electric light in	
opal globe	2.00 ..
Ditto ditto, frosted globe.. ..	3.50 ..

We have therefore, at last, a series of figures on this important subject which cannot possibly be doubted. The experiments which have yielded them have been carried out with the greatest care, and every precaution has been taken to render them so correct that all future cavil or controversy will be avoided; the practical outcome of the matter being that, used in the best possible manner, the amount of illumination in both cases being equal, the electric light costs $1\frac{1}{2}$ times as much as gas.

PROCEEDINGS OF SOCIETIES.

PHYSICAL SOCIETY.

Ordinary Meeting, May 24, 1879.

Prof. W. G. ADAMS, President, in the Chair.

New members:—Mr. R. Sedley Taylor, M.A., and Mr. Walter Emmott, A.S.T.E.

Mr. W. J. WILSON exhibited a new harmonograph and figures drawn by it. The figures drawn by prior harmonographs are all more or less imperfect, owing to loss of motion in the pendulums actuating the marking pen; and Mr. Wilson therefore designed a new harmonograph, which not only gives perfect figures, but admits of the phase of either of the two compounded motions being decreased by a known amount. In this instrument toothed wheels take the place of pendulums, the ratio of the teeth giving the ratio of the periods of the motions. By employing the device of an intermediate wheel gearing with two others, and arranging two or more wheels on the intermediate axle, a great variety of phase can be

obtained for each motion. An ingenious adjustment by means of a movable nut allows the phase of either or both motions to be altered to a known extent, and thus an endless variety of figures can be obtained. As in other harmonographs, a writing table, on which the paper is placed, and an aniline glass pen, are used. Several of the figures done also on glass were thrown on the screen, the stereoscopic effect being very apparent. In reply to a query, Mr. Wilson said that he had adapted some of the figures to the stereoscope.

Prof. HUGHES explained his new Induction Balance, and showed some of its principal effects. It is well known that on an electric current passing along a wire adjacent to another wire, an induced current is set up in the second wire in an opposite direction to the first or primary current. In the induction balance two primary circuits or coils are taken, with the same current (interrupted by a microphone acted upon by the ticking of a clock) running through both; and between these is placed a secondary circuit or coil in connection with a telephone. The primary coils are so wound as to oppose each others induction on the intermediate secondary. There is a point, moreover, between where these opposed inductive influences exactly neutralise each other. If the secondary coil be placed there, no induced ticking of the clock can be heard; but if the secondary be displaced to one or other side of this point, the ticking can be heard in the telephone, increasing in loudness as the secondary approaches one or other of the primaries. If the distance between the primaries be graduated into a scale, this contrivance becomes a sonometer, since it gives an absolute zero of sound and degrees of loudness. It is well adapted for measuring the hearing power of the ear, and when used for this purpose it is known as the audiometer. By splitting the secondary coil into two parts and placing each close to its proper primary, so that there are four coils in all arranged in two pairs, the extremes in one primary circuit and the means in one secondary, the two opposing parts of the balance can be separated from each other, so as not to disturb each others action, and the balance made very sensitive by the closeness of the primaries and secondaries. Prof. Hughes finds that there is a line or zone of maximum induction midway between each primary and its secondary. If a conductor, such as a piece of metal, be put in this position it has a maximum distributing effect on the balance, due probably to the electric currents generated in it by the induction. The effect is apparently proportional to the conductivity of the metal. It requires an exactly similar piece of metal put between the other pair of coils to restore the equilibrium of the balance. A very slight difference of alloy or of weight between two like coins is at once observed from the imperfect restitution of the balance; base coins are thus also at once detected. Again, it is possible for a person to tell what particular coin or coins are in one part of the balance by trial of the same coins in his part. When plates of non-magnetic metals are held vertical in the balance their distributing effect is nil; iron, on the other hand, gives its maximum effect at this position, because its maximum effect overbalances its electrical effect. Two pieces of iron may therefore neutralise each other as cores in an induction coil. Steel is easy to balance compared with soft iron. Zinc disturbs most when placed along the sides of each pair of coils; iron when in centre. A certain length of metal laid along the outsides of the coils produces silence. The maximum line of inductive force is midway between the coils, and there is a line of minimum force at half the thickness of each coil; a metal placed at these lines of minimum force has no disturbing effect on the balance. Pressure, torsion, heat, magnetism, strain, and in fact all imponderable forces, are indicated, and their effects may be measured.

Prof. W. G. ADAMS believed that one result of Prof. Hughes experiments will be the determination of the way in which the law of electro-dynamic induction depends on density. He also imagined that the metal

the maximum line between the coils gathered the lines of force to itself, whereas when on the minimum lines it could not thus divert them.

Prof. AYRTON cited the early experiment of Faraday with a sheet of copper oscillating to rest between two opposite magnetic poles. The copper took a long time to stop; but a sheet of iron placed between two like poles was soon stopped, owing to its becoming imbued with an opposite polarity, and deflecting the lines of force. He also suggested that the divergence of the results for conductivity of metals got by the induction balance from those got by the Wheatstone balance might be due to that electric inertia suspected by Sir William Thomson.

Prof. GUTHRIE thought that the induction balance pointed to the conclusion that the disturbing effect of a conducting mass applied in this way is proportional to the quantity of electricity generated in it under certain conditions of temperature, &c. The determination of the conductivity of liquids would be a useful application of the balance.

Mr. CHANDLER ROBERTS gave some results which he had obtained from an examination of certain alloys by means of the induction balance. He had been able to detect a difference of 1 part in 1000 in the amount of silver present in two shillings of equal weight. He also pointed out that Mathiessen divided alloys into three classes:—(1) Solidified solutions of one metal in another; (2) Solidified solutions of one metal in an allotropic modification of another metal; (3) Solidified solutions of allotropic modifications of both metals. For the first class the curve of electric conductivity is a straight line; for the second, a parabolic curve; for the third, a bent line. Mr. Roberts found that the balance gave the characteristic curve for the first class with an alloy of lead and tin, and for the second with an alloy of gold and silver. With a copper-tin alloy, which is a good example of the third class, he found the curve given by the balance to be intermediate between Alfred Rich's curve of density and Mathiessen's curve of conductivity, and considers that the balance is influenced by the density as well as by the conductivity of the metal interposed.

Prof. HUGHES said that as the working of metals appeared to effect their conductivity, he was inclined to rely more on the conductivity measurements of the balance than on those of the Wheatstone bridge. By the balance plain pieces of metal were taken, whereas by the bridge wires were mostly taken. He would rather not give any theories yet as to the results obtained from the balance.

Dr. ERCK then exhibited his novel Pump for lifting solutions out of batteries.

chemists, more remote. Besides this, the difficulty of hunting down a practical fact amongst a crowd of theories is avoided.

In the present issue the author has adhered more or less closely to the lines followed in the first edition, which was published in 1874. As then, he divides the elements and pseudo-elements into monads, dyads, &c.—an arrangement which greatly helps the memory. The chapter on Chemical Theory has also undergone the process of subdivision, and the matter has been greatly extended. The chapters on the Organic Compounds have also received large additions, and have also been relegated to the concluding chapters of the book instead of being classified according to their chemical constitution as in the first edition. This arrangement, though perhaps not very philosophical, is at any rate a convenient one for the hard worked medical or pharmaceutical student.

Two useful appendices have been added to the new edition: one showing the quantities of the various ingredients to be added in parts by weight to 100 parts of alcohol, to produce the different artificial fruit essences; the other, the relation between the density and flashing point of paraffin oils. These appendices will be of great use to the practical chemist, but why annex them to the theoretical division of the work?

The second volume of the work has been in a great measure remodelled and re-written. The processes employed by chemists are first described, beginning with solution and lixiviation, and ending with electrolysis and pyrology. The next chapter treats of the detection and separation of the metals used in pharmacy, and Chapter 3 follows with the detection and separation of the acidulous radicals. Chapter 4 gives the method of quantitative analysis as applied to the examination of the official salts—*official*, by the way, being the new and more correct epithet applied to what used to be called *official* preparations. The next chapter gives a course of quantitative analysis as applied to the detection of unknown salts, while Chapter 6 gives a complete course of alkaloidal analysis. The other chapters are devoted to toxicological analysis, volumetric and geometric quantitative analysis in general, ultimate organic analysis, and, lastly, special analytical processes, such as the analysis of potable water and cinchona bark, the valuation of opium, the estimation of emetine in ipecacuanha, the alcoholic strength of spirits, wines, and tinctures, and the analyses of urine and urinary calculi.

Mr. Joseph Ince has again done good service to Dr. Muter by undertaking the tedious—but, as far as his students go, certainly not the thankless—task of compiling copious indexes to both volumes, which will greatly add to the value of Dr. Muter's already valuable work.

NOTICES OF BOOKS.

An Introduction to Pharmaceutical and Medical Chemistry (Theoretical and Descriptive), arranged on the principle of the Course of Lectures as delivered at the South London School of Pharmacy. Second Edition. By Dr. JOHN MUTER, M.A., F.C.S., &c. London: Simpkin and Marshall. 1879.

An Introduction to Analytical Chemistry, &c. Second Edition. By Dr. JOHN MUTER, M.A., F.C.S., &c. London: Simpkin and Marshall. 1879.

Dr. MUTER has done well to divide the second edition of his "Pharmaceutical and Medical Chemistry" into two separate volumes. The first part, being almost purely theoretical in its character, is intended to be for perusal at home, while the second half, being purely practical, is to be used in the laboratory. Besides obviating the inconvenience of dragging a heavy book backwards and forwards from the study to the laboratory, the division of the Manual into two parts will render the possibility of spoiling an expensive work, by the accidents common to all

Report on the Air of Glasgow, chiefly relative to Enclosed Spaces and Smoke. By W. J. DUNNACHIE, in co-operation with the Medical Officer of Health. Presented to the Committee of Health of the Magistrates and Council of Glasgow. May, 1879.

It is found that the proportion of atmospheric sulphur reaches its annual minimum in July, and, as might be expected, is highest in the central stations. Nitrogen, in the forms of actual and potential (albuminoid) ammonia, has two periods of excess, one in the summer and one in the winter. It appears, from Mr. Dunnachie's experiments, that these two forms of ammonia are largely derived from the combustion of coal, &c., and are consequently no absolute indices of the sanitary condition of the air. This fact has scarcely received sufficient attention on the part of sanitary chemists. Even Dr. R. A. Smith, in his classical work on "Air and Rain," though fully admitting the presence of tarry distillates in the atmosphere, still says "the ammonia is one measure of sewage of air not purified." Dr. J. B. Russell observes that "the methods of Pasteur and Pouchet, of Cohn and Lister, are necessary to give precise meaning to the work of Angus Smith

and other chemists in the region of air, and of Wanklyn and Frankland in the region of water." The guidance of chemical analysis is not absolute and all-sufficient; but as the most dangerous impurities, whether of air or of water, appear to be not merely organic, but organised, it must be controlled and supplemented by biological inquiry. This, however, neither Dr. Angus Smith nor Prof. Frankland will, we think, dispute.

Proposed Legislation on the Adulteration of Food and Medicine. By E. R. SQUIBB, of Brooklyn. New York: G. B. Putnam's Sons.

THIS pamphlet comprises the draft of an anti-adulteration law for the State of New York. It includes definitions of food, of medicine, and of the offence of adulteration, the various phases of which are illustrated by examples. The penalty for a first offence is a fine not exceeding two hundred dollars, and for further convictions imprisonment with hard labour for a term not exceeding six months. The Act is to be worked by a State Board, composed of two physicians, one chemist and physicist, one lawyer, and one "business man."

The author gives an abstract of English legislation on the adulteration question, and a history of their working. Here he speaks of analysts who "took the initiative, bought their own samples for analysis, and entered upon prosecution where necessary." This is a misapprehension. A public analyst who should act in this manner would be gravely overstepping his duty.

CORRESPONDENCE.

MANUFACTURE OF SULPHURIC ACID.

To the Editor of the Chemical News.

SIR,—I was pleased to note in the *CHEMICAL NEWS*, vol. xxxix., p. 226, only just to hand, that Dr. Lunge qualifies the favourable opinion on the use of liquid nitric acid, given on his authority in Lock's work, lately published, on the manufacture of sulphuric acid; for, I confess, I was somewhat surprised on reading the book, to find that, with his extended practical experience, he should advocate a process which entails expenses in the shape of labour, fuel, and maintenance of extra plant, as well as unavoidable loss and waste. After undertaking the management of these works I seized the first opportunity of setting up two large Daglish's nitre pots, and to discard the nitric acid stills. The result is a saving of 1½ per cent of saltpetre on the 100 tons pyrites burnt weekly. The requisite quantity of nitre is weighed out twice daily to the chamber foreman, and is proportionately charged into the pots once every hour.—I am, &c.,

JOHN COX.

Chemische Fabrik "Colombia,"
Mülheim-a-Rhein, May 26, 1879.

EXPLOSIONS IN FLOUR-MILLS.

To the Editor of the Chemical News.

SIR,—Will you allow me, in reply to Mr. Arthur M'Dougall's letter, briefly to state that my letter on the above subject appeared in the *Glasgow Herald* immediately after the explosion in the Tradeston Flour Mills; in fact, I believe, about a couple of days after the sad accident. It was noticed in a brief leading article on the subject as the most likely solution of the mystery, and after Dr. Stephenson Macadam's paper appeared (which was considerably afterwards), the Scottish Royal Society appointed a commission of inquiry into the subject, together with the question of priority, the upshot of which was

that I received a private letter from the Secretary, Dr. Ferguson, of Edinburgh, which I still have, in England), in which my prior claims were fully acknowledged, and I was informed, too, that my report or letter to the *Glasgow Herald* was framed as the first. Dr. Macadam was mentioned, too, as having been the first to invite a scientific investigation to the theory I brought forward, and so to support it further. Still, I must conscientiously acknowledge that the brain which to the best of my knowledge deserves the fullest credit of the conception is that of the Austrian chemist, whose name I have forgotten, but whose little notice in *Dingler's Journals* supplied me with the necessary light on the subject, which I did but endeavour to spread and throw upon that fatal catastrophe at Glasgow. Mr. M'Dougall's letter will be welcomed as a very interesting contribution, and I feel no doubt he is right as to the abolition of the stove-room. With regard to the inconvenience of small electric lights disposed over a flour-mill, I cannot judge, not having experience in the matter. It is for the mill-owner to decide which inconvenience is greater, that of ordinary lights, with the chance of explosions; or the electric lights, with safety. I fancy the filtration of the air, as Mr. M'Dougall suggests, may prove a difficult matter; but I wish it all success, as it would doubtless remove the danger. Still, it is always to be remembered that from abnormal causes and accidents, the air may sometimes even then become laden with flour dust in certain rooms or parts of the mill, and a vivid flash might communicate itself where an ordinary flame or light would fail to do so.—I am, &c.,

WATSON SMITH, F.C.S., F.I.C.

Zürich, June 1, 1879.

DR. PAVY'S METHOD OF DETERMINING GLUCOSE.

To the Editor of the Chemical News.

SIR,—It was only a few days ago that my attention was drawn to the article by Mr. Otto Hehner in the *CHEMICAL NEWS* (vol. xxxix., p. 197) on the ammoniated cupric liquid proposed by me for the quantitative determination of sugar. I am glad to find that so able a chemist as Mr. Hehner has conducted an inquiry into the merits of the test, and it is satisfactory to notice that with a proviso regarding the amount of alkali employed he is able to speak of the method as "capable of furnishing excellent results."

At the outset of his investigations Mr. Hehner obtained results which stood at variance with mine, and considerably paid a visit to my laboratory at Guy's Hospital to see in what manner the discordancy was to be accounted for. Upon being informed of his mode of procedure, which consisted in using 100 c.c. of the test-liquid, and dropping the saccharine solution in slowly throughout the process, instead of employing only 20 or 40 c.c., and running the liquid in rapidly from the burette until near the point required, and then proceeding slowly—a difference which leads to the operation being much more prolonged in the one case than in the other, it immediately occurred to me that the discordancy arose from the test-liquid failing to resist the influence of the prolonged boiling, and that this would probably be rectified by the employment of a larger amount of alkali. I have found with Fehling's solution that its stability under ebullition is impaired by dilution unless an additional quantity of alkali is put into it. I suggested to my assistant, Mr. Scard, who has ably worked under my direction in this matter, that more alkali should be used, and it was found that the true explanation of the discordancy had been hit upon. Without further concert than I have mentioned, the idea also occurred to Mr. Hehner to try the effect of a larger quantity of alkali, and when an interview took place a short time afterwards each learnt that an accord had been arrived at.

In my communication to the Royal Society I stated that with the test proposed, 1 atom of sugar appropriated

6 atoms of oxide of copper instead of 5 as is the case with Fehling's solution used in the ordinary way. I further mentioned that potash might be added to the extent of 1 grm. to 20 c.c. of the ammoniated test without altering the result, but that with the addition of 5 grms., or anything beyond, 5 instead of 6 atoms of oxide of copper were appropriated by 1 atom of sugar, and with quantities of added potash between the 1 and 5 grms. the results stood between the 5 and 6 atoms of oxide of copper. Mr. Hehner, in commenting upon this, says that his observations, which he has verified over and over again, show that a much smaller excess of soda influences the result than is stated by me. Mr. Hehner seems not to have noticed that the added alkali in my observations consisted of potash instead of soda, which he employed. The difference in the atomic weights of the two alkalies will nearly suffice to account for the difference existing in our recorded results.

For many years past the liquid which has been used in my laboratory as a sugar-test has been made with potash instead of soda, and has contained nearly double the equivalent of the soda existing in Fehling's solution. I was led to introduce the larger amount of alkali because it appeared to me that the liquid possessed greater stability and precision of behaviour. I am now using this liquid not only as formerly, but also for the preparation of the ammoniated solution, and it contains a sufficient amount of alkali to serve for what is required. The quantity of copper is the same as in Fehling's solution. The two liquids therefore possess the same quantitative value. The composition is as follows:—

Cupric sulphate.	34.65 grms.
Potassio-sodic tartrate (Rochelle salt)	173 "
Potash	160 "
Water to 1 litre.	

For the ammoniated test 120 c.c. of this solution are mixed with 300 c.c. of strong ammonia (sp. gr. 0.880), and water is added so as to make up to a litre. The test, being in this form a 6-atom instead of a 5-atom oxidising liquid, it is of 1-20th the value in relation to sugar indication of the original solution, or 20 c.c. correspond with 0.010 grm. of glucose.

On account of the liability of the cupric sulphate to impurity it is preferable, where great accuracy is required, to use electrolytic copper. The quantity of copper to be taken is 8.808 grms. This is dissolved in nitric acid and evaporated to dryness after the addition of sufficient sulphuric acid to secure that a sulphate is left. The product is then dissolved in water and neutralised with potash. In this way the exact amount of cupric sulphate is supplied that is required to yield a solution of the standard strength.

I originally recommended, for preventing the atmosphere from becoming charged with the evolved ammonia, that a U-shaped tube containing fragments of moistened pumice-stone should be used. I find that a much more simple and convenient contrivance for attaining the object is afforded by carrying a piece of vulcanised tubing from the flask into a beaker or other vessel containing water. The end of the vulcanised tubing intended to dip into the water is plugged at the extremity, and just above the plug provided with a transverse slit through three-fourths of its extent. This admits of the escape of air and ammoniacal vapour from the flask, but precludes by its valvular arrangement the sucking back of water from the occurrence of any sudden condensation in the interior of the flask.

Uric acid exerts a reducing action upon oxides of copper equally as precise as glucose. The quantitative determination of it has hitherto constituted a lengthy and not very satisfactory process, but I find that with the ammoniated cupric liquid its determination may be easily and speedily effected. A large number of observations have been conducted in my laboratory with the view of ascertaining its precise reducing capacity, and the results so closely conform with the expression that 3 atoms of oxide of copper are reduced by 1 atom of uric acid that I am led

to give this as the relation existing. I am not aware that the extent of reducing action of uric acid upon the oxide of copper has hitherto received attention, and I should be glad to learn if the conclusion I have mentioned is confirmed by the observations of others.—I am, &c.,
F. W. PAVY.

May 30, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 17, April 28, 1879.

The Electric Light.—J. Jamin.—The burner submitted to the Academy with its points downwards has considerable advantages; such as simplicity, since it requires no mechanism and requires no preliminary preparation, beyond a support and the coke points; mechanical economy, since the number of flames is almost doubled; augmentation of light, since each of the new foci is almost twice as powerful as those of the old construction; quality of the light, which is whiter; more advantageous arrangement of the foci, which direct their greatest quantity of light downwards, where it is wanted, instead of up into the air, where it is useless; and, lastly, economy of combustible material.

Electric Inscription of Words.—M. Boudet de Paris.—The transmitting apparatus is a microphonic speaker, the carbons of which, instead of being pressed by a spring, are simply maintained in contact by the pressure of a small piece of paper folded in form of a V. The vibrations of the diaphragm of the receiving apparatus cannot be written, since the movements of the style, however delicate the apparatus, can scarcely be distinguished upon the lamp-black. To enlarge the magnetic vibrations of the receiver the cover and the diaphragm of a Bell's telephone are taken away, and on the wood of the instrument there is fixed the end of a small stiff steel spring. The other end of the spring abuts on the surface of the magnetic nucleus surrounded by its coil; to this extremity is soldered a small mass of soft iron, weighing about 10 grms., and upon this mass and in the produced line of the axis of the spring is fixed a light style of bamboo, 10 centimetres in length and terminating in a slender whale-bone pen.

Theoretic and Experimental Demonstration of the following Definition of Temperature; the Temperature is Represented by the Length of the Calorific Oscillation of the Molecules of a Body.—R. Pictet.—A mathematical paper, not capable of useful abstraction.

A Siren with an Electro-magnetic Regulator.—M. Bourbouze.—By means of this instrument sound can be made to pass from 8162 vibrations per second, through all the intermediary notes to 128 vibrations.

The Laws of Dissociation.—MM. Moitessier and R. Engel.—The dissociation of a body both whose components are volatile takes place even when the body is placed in presence of one of the products of dissociation, so long as the tension of this product does not surpass the dissociation-tension of the body at the temperature of the operation. When the tension of one of the components is greater than the dissociation-tension of the compound the dissociation no longer takes place. Two cases may then arise; either the dissociable compound is volatile, and in that case the true vapour density may be determined, or else the dissociable body is not volatile, as is the case with chloral hydrate at 60°. When two gaseous products on combining form a dissociable com-

pound the combination only takes place when the sum of the tensions of the components exceeds the dissociation-tension of the compound.

Determination of Glucose in Blood.—P. Cazeneuve.—A reply to the criticisms of MM. d'Arsonval and Picard (*Comptes Rendus*, 1879, pp. 753 and 755).

Contributions towards the History of Beer-yeast and of Alcoholic Fermentation: Physical and Physiological Action of Certain Substances, Saline or otherwise, upon Normal Yeast.—A. Béchamp.—Yeast which has undergone the influence of sodic acetate, and which has been well washed and drained, can no longer be as completely fluidified by a new addition of acetate. Yeast, after a first treatment with the acetate, and whilst still impregnated with this salt, is still capable of undergoing fermentation. Yeast which has been, after two or three treatments with acetate, deprived of the greater part of its soluble matters is nevertheless capable of setting up energetic fermentation in cane-sugar.

No. 18, May 5, 1879.

Certain Derivatives of Durol (α -Tetra-methylbenzol).—MM. Friedel, Crafts, and Ador.—The authors have obtained a compound, which may be named phenyl-duryl-carbonyl, if we give the name duryl to the monatomic residue of durol, from which 1 atom of the hydrogen of the benzenic nucleus has been removed. At the same time there is formed another compound, almost insoluble in boiling alcohol, duren-dicarbonyl-diphenyl or duren-dibenzoyl, which, if treated at a boiling heat with melting potassa, is resolved into benzoic acid and durol.

Crystals extracted from Cast-iron by means of Ether or Petroleum.—J. Lawrence Smith.—On treating comminuted cast-iron with these solvents, on the spontaneous evaporation of the liquid, the author obtained acicular crystals consisting principally of sulphur, and absolutely similar to those extracted by the same process from meteoric iron. M. Berthelot, to whom the author had forwarded specimens of the crystals, stated that he had obtained similar bodies by the action of pure ether upon octahedral sulphur and upon anhydrous iron sulphides. It thus appears that even neutral solvents do not in all cases act by simple solution upon the bodies with which they come in contact, but effect also a chemical alteration.

Effects of Carbonic Sulphide upon the Radicular System of the Vine.—M. Boiteau.—The author concludes that this agent in doses of 6 to 10 grms. destroys by poisoning all the parts of the root system which are within a radius of about 10 centimetres of the point where it is applied. This action is produced in the parts situate from 20 to 35 centims. below the surface. He still recommends the use of this sulphide, applied at a distance of more than 10 centims. from the main root.

Thermic Formation of Hydrogen Silicide.—J. Ogier.—The heat disengaged in the combustion of 1 equiv. of this compound is 324.3 cals., and the mean heat of its formation is 24.8 cals.

Limit of the Separation of Alcohol and Water by Distillation.—J. A. Le Bel.—The greatest degree of concentration obtained by repeated rectifications was 96.5 per cent. On rectification over quick-lime an alcohol of 98.5 was produced. When this spirit was submitted to fractional distillation the water passed over first, and the residue was the most highly alcoholic. After three rectifications the first portions marked 97.4 and the residue 99.3. The author finds that amylic alcohol from wines has not the repulsive odour of fusel oil or of the crude amylic alcohols of (beet-root) treacle. This odour, and possibly the injurious physiological action of the higher alcohols, may possibly, he considers, be due to the presence of certain empyreumatic compounds.

A New Isomer of Angelic Acid.—E. Du villier.—

The author has obtained iso-angelic acid along with ethyl-oxy-valeric acid on causing the bromo-iso-valerate of ethyl to act upon the ethylate of sodium in an alcoholic solution.

Transformation of Camphic Acid into Camphor.—J. de Montgolfier.—On heating calcic camphate and formates together the author obtained camphor, well characterised by its composition and properties, and accompanied by camphren, an isomer of phoron.

No. 19, May 12, 1879.

The Vision of Colours, especially the Influence exercised upon the Vision of Coloured Objects in Rotatory Motion when Observed Comparatively with Identical Bodies in a state of Rest.—E. Chevreul.—This is a somewhat diffuse extract from a recent work by the author, and does not admit of useful abstraction.

Bases Derived from Aldol-ammonia.—A. Wurtz.—Ammonia, when reacting upon aldol, gives rise to various bases, whose nature varies according to the conditions of the experiment. At 100° there are formed certain oxygenous bases soluble in water, the study of which is still incomplete. At 140° to 180°, and in presence of an excess of ammonia, dark-coloured oily bodies are formed, soluble in ether. A mixture of liquid and volatile bases is obtained on submitting aldol-ammonia to dry distillation in a current of ammoniacal gas.

Effects of Inhaling Oil of Turpentine.—M. Poincaré.—The author has examined 282 workmen who use this oil in their trades, and has kept animals for several months in a medium strongly charged with its vapour. Among the workmen the symptoms were headache, dizziness, watery eyes, weakness of sight, especially in artificial light, cough, and troubled digestion. In most cases the constitution became habituated to this agent, but sometimes a change of employment was necessary. The animals experimented upon remained in a normal condition if good ventilation was maintained. In confined air death ensued in consequence of congestion of the brain, lungs, &c.

Two Applications of the Method of MM. Fizeau and Foucault.—M. Mouton.—A mathematical paper not suitable for abstraction.

Thermic Researches upon Silicic Ether.—J. Ogier.—The mean heat of formation of silicic ether is -11.5 cals.

Action of Ammoniacal Salts upon Certain Metallic Sulphides, and Application of the Facts Observed to Mineral Analysis.—Ph. de Clermont.—Bismuth, cadmium, and copper sulphides are not affected if boiled with a solution of sal-ammoniac, as is also the case with mercurous and mercuric sulphides. Antimony trisulphide is completely decomposed by sal-ammoniac, yielding antimony sulphide, which escapes, and antimony chloride, which remains in solution. Stannic sulphide yields stannic acid and no tin dissolves. Stannous sulphide is similarly decomposed, leaving stannous oxide. Metals not thrown down by hydric sulphide from their acid solutions, but converted by ammoniac hydrosulphate either into sulphides or insoluble oxides, after the action of this reagent, behave in a peculiar manner with sal-ammoniac. It is already known that manganese sulphide dissolves as chloride. Iron sulphide in the same manner gives iron chloride. Nickel and cobalt sulphides dissolve likewise, though more slowly. Zinc sulphide resists longer, but ultimately dissolves also. Alumina and chromic oxide precipitated by ammoniac hydrosulphate are known to be insoluble in sal-ammoniac. On these reactions the author founds a method for the separation of certain metals. If a solution contains cobalt, nickel, manganese, iron, alumina, chrome, and zinc, it is precipitated with ammoniac hydrosulphate, the mixture added to a boiling solution of sal-ammoniac, and kept in ebullition for a sufficient time. On filtering, which is effected rapidly, the filtrate contains all the iron and manganese, part of

the cobalt, nickel, and zinc, whilst the undissolved portion contains all the alumina and chrome with the residue of the nickel, cobalt, and zinc. The analysis is completed by known methods. If iron and manganese alone have to be separated from alumina and chrome the process is complete and precise.

New Formation of Glycocoll by means of Nitro-acetic Ether.—M. de Forcrand.—Nitro-acetic ether yields glycocoll on reduction with tin and hydrochloric acid.

No. 20, May 19, 1879.

A New Derivative of Nicotin.—A. Cahours and A. Etard.—The new base, to which the authors assigned the name thio-tetra-pyridin, has a composition represented by $C_{16}H_{18}N_4S_2$.

Transparence of the Media of the Eye for the Ultra-violet Rays.—J. L. Soret.—The author concludes that the absorption of the totality of the media of the eye must render it impossible to perceive rays whose refrangibility exceeds that of the extreme radiations of the solar spectrum, say the ray U.

Preliminary Researches on the Action of Acids upon Salts without the Intervention of a Solvent.—M. Lorin.—The direction of the results indicates chemical action more or less distinct, and which, for the fatty acids, decreases from formic acid to each of its successive homologues.

Presence of Mercury in the Mineral Waters of St. Neaire.—E. Willm.—The author's results have been mutually contradictory. Supposing mercury to be a constant element of the water the quantity shown by his only affirmative experiment falls far short of that resulting from the analysis by Dr. Garrigon.

The Mode of Combination of Iron in Hæmatoglobulin.—L. Jolly.—The author concludes from his analyses that iron occurs in the blood-globules merely as phosphate.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 64, April, 1879.

Report presented by M. Ed. Collignon on the Electric Brake of M. A. Achard.—The moment that an electric circuit is broken, whether by the separation of a part of the train, by carriages getting off the line, or by fire, the wheels are automatically locked.

Report by M. Sebert on the Electric Bells of M. Laville.—The arrangement of this apparatus cannot be made intelligible without the aid of the accompanying illustration.

Revue Universelle des Mines, de la Metallurgie, &c.,
Tome 4, No. 3, November and December, 1878.

Explosions Caused by Dust.—This paper recounts the facts laid before the Société d'Encouragement by Prof. Lawrence Smith in November, 1878, and the discussion on the same subject at the Academy of Sciences.

No. 1, January and February, 1879.

Relation between the Temperature of the Earth and the Depth below the Surface.—P. Van Dijk.—The author draws the following conclusions:—The increase of the geothermic scale proves that the earth is engaged in a continual process of cooling. This increase corresponds to the law of the refrigeration of a spherical body, having a uniform elevated initial temperature. The degree of cooling differs much for different parts of the earth's surface, according to their geographical and thermological position, and exercises an important influence on

the vertical distribution of underground heat in any place. The loss of heat which the earth incessantly undergoes, from its surface being infinitely small, compared with the quantity stored up within, a large part of the earth's interior must still possess a very high temperature. The real geothermic progression, as accessible to human observation, far from indicating a minimum thickness for the earth's crust, imposes, on the contrary, a very narrow limit on the maximum temperature of the earth's interior. By the abstract application of this progression we should be led to ascribe to the earth's centre a heat not exceeding 100° . Actual facts observed, however, cannot be more simply explained than by the reaction of an igneous or fused nucleus, of a crust of considerable thickness, and having little conductivity for heat. The thickness of the earth's crust is very unequal, probably three times greater in the polar regions than in the volcanic regions of the Equator, and in one and the same latitude greater beneath the sea than on dry land. The greatest differences of thickness for one and the same latitude are met with in countries where, with an elevated mean temperature at the surface, the alternation of dry land and deep sea is most frequent. The volcanic character of the Eastern Archipelago agrees with this conclusion. According where the terrestrial crust is thinnest a temperature of 3000° will be found at the depth of 21 geographical leagues, not more than 1-40th of the earth's radius. In the polar regions it is probably more than four times as thick.

Biedermann's Central-blatt.

Heft 3, March, 1879.

Preparation of Artificial Animal Charcoal.—Th. Filter.—The author proposes to obtain a cheap form of gelatin by treating leather waste mixed with from 1 to 5 per cent of caustic soda with steam at a pressure of six atmospheres in closed iron vessels. The resulting semifluid mass is placed in a centrifugal, which keeps back the gelatin whilst tannate of soda runs off. So much of this jelly as is equal to 33 kilos. of dry glue is mixed with 50 kilos. bibasic phosphate of lime and 17 kilos. phosphate of magnesia. The mass is submitted to strong hydraulic pressure between layers of felt, dried at 110° , and ignited in the ordinary manner.

Occurrence and Life of Bacteria and their Influence upon Yeast.—MM. Schnetzler, Bretefeld, Downes, Blunt, and Gunning.—Light is hostile to the development of bacteria and microscopic fungi. The exclusion of oxygen occasions the death of bacteria, but promotes putrefaction.

Reimann's Färber Zeitung,
No. 16, 1879.

Tartar Emetic in Dyeing.—The author points out that tartar emetic, as used in cotton-dyeing, serves not to fix the aniline colours themselves, but merely to fasten the tannin, thus playing the part of an indirect mordant. Several experiments have been instituted at the Färber Akademie in order to ascertain whether this application of antimony can be pronounced injurious to public health. It was found that water, in which cotton yarns dyed with aniline colours on a mordant of tannin and tartar emetic had been steeped, or especially boiled, gave distinct indications of antimony when tested in ordinary manners. The quantity of the metallic compound fixed on the fibre seems, indeed, far too trifling to have any effect upon human health. Still, in view of the panic which has seized upon the public mind, and of the tendency of the literary and political press to attribute all mysterious attacks of sickness to the influence of poisonous dyes, &c., Dr. Reimann—in our opinion very judiciously—advises dyers to dispense with the use of antimony, which will doubtless be found by no means necessary.

No. 17, 1879.

This issue is mainly taken up with the specification of F. V. Kallab's patent process for bleaching animal fibre with hydrosulphurous acid.

Chemiker Zeitung.

No. 16, April 17, 1879.

Inflammability of Petroleum.—The first part of a paper on the possible sources of error in testing petroleum, and the precautions needful for obtaining comparable results.

Poisonous Colours.—The Association for the Protection of Chemical Industry, in Germany, has petitioned the Reichstag that no colour shall be condemned as dangerous until it has been duly examined by a Commission of specially qualified chemists and physiologists.

Dr. Volhardt succeeds to the chair of the late Prof. Gorup-Besanez, at the University of Erlangen.

Properties and Chemical Examination of Tea.—Dr. J. M. Eder.—The author holds that one-half of all tea sold is sophisticated with spent tea-leaves. He places no confidence in the determination of thein as a test of the genuine character of tea, but proposes to ascertain total extractive matter as obtained with hot water; proportion of tannin in the decoction; total ash; proportion of ash soluble in water.

Starch in Filter-paper.—The *Beyer Industrie-Blatt* cautions chemists against errors which may be produced by this impurity.

Gold in Petroleum.—According to the *Bergmann* Mr. J. Turnbridge has extracted 34 dollars' worth of gold from a ton of the residues from petroleum-stills. The source of the petroleum was not ascertained.

Salubrine-Cenosote.—This name has been employed by M. Perrot, of Geneva, to a mixture of salicylic acid and tartar, which he recommends as a preservative for beer, wine, &c.

A Curiosity of Patent Literature.—A specific for the teeth, patented in Belgium, consists of 950 parts water, 12 parts urea, 0.4 part uric acid, 7 parts of fixed salts, and 8.90 of urate, oxalurate, oxalate of lime, cystin, &c.—a composition which agrees wonderfully well with that of urine. A little perfume has been added. (Is this a new way out of the sewage difficulty?)

No. 18, May 1, 1879.

According to the *Repert. de Pharmacie* magenta has been successfully administered in the treatment of albuminuria, to the extent of half a gramme daily.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 15, April 10, 1879.

Origin of Lactose.—There appears to exist in the mammary glands during lactation a lactogenous substance, which, however, differs widely from the other glycogens. M. Schützenberger is engaged with its chemical examination.

Quantitative Analysis of White Light.—Prof. Rood divides the spectrum into twelve parts, and, multiplying the space occupied by each part by the relative luminous intensity, he obtains the following numbers:—Red, 54; orange-red, 140; pure orange, 80; orange-yellow, 114; yellow, 54; greenish yellow, 206; yellowish green, 121; green and greenish blue, 134; Prussian blue, 32; blue, 40; violet, approaching to ultramarine, 20; pure violet, 5. Hence the quantity of light in the so-called warm colours is three times greater than that in the cold colours. This division of the spectrum does not seem to rest upon any demonstrable basis, and the author has not been guided by the spectral lines.

No. 16, April 17, 1879.

Reduction of the Cost of the Electric Light.—M. Delaurier.—By preventing all access of atmospheric air the duration of the carbon points is increased ten-fold, and the working cost is reduced by two-thirds.

Physiological Action of Benzol.—M. Bensch.—The author shows that benzol, which has been proposed as an anæsthetic, acts as a poison if inhaled as vapour or introduced into the circulation.

Spectroscopic Optics.—M. Moigno, in translating Prof. Piazzzi Smyth's lecture on "End-on Illumination," points out that he has been to some extent anticipated by Dr. van Monckhoven (*Les Mondes*, 1877, p. 177).

No. 1, May 1, 1879.

This issue contains no original chemical or physical matter.

No. 2, May 8, 1879.

Universal Support, or Electro-diapason for Representing Vibratory Movements and showing them in Projection.—A. Dubosc.—This memoir cannot be intelligibly abstracted without the addition of the five accompanying plates.

No. 3, May 15, 1879.

An international subscription is proposed to erect a monument to Nicéphore Niepce, at his native town, Chalon-sur-Saône.

The new freezing machine of M. D. L. Holden, of Philadelphia, is, in sum, that of M. Piçet, somewhat varied in form.

The electric bit is applied very successfully to vicious horses by the Omnibus Company of Paris.

MISCELLANEOUS.

Mineralogical Society of Great Britain and Ireland.—A general meeting was held at 116, Victoria Street, London, S.W. (Prof. P. G. Bonney in the chair), on Tuesday, June 3rd. The election of fourteen corresponding members, eight ordinary members, and two associates was announced. The following papers were read:—"On Abriachanite, a new Scottish Mineral," by Prof. M. F. Heddle and Dr. W. Aitken; "On Haughtonite, a new Mica," by Prof. M. F. Heddle; "On Bheekite and Xantholite from Scotland," by Prof. M. F. Heddle; "On Christophite from St. Agnes, Cornwall," by J. H. Collins, F.G.S.; "Minerals from Japan," by John Milne, LL.D.; "On some Gold Occurrences," by the Rev. J. Clifton Ward, F.G.S.; "Additional Note on Penwithite," by J. H. Collins, F.G.S.; "Measurements of the Angles of Basaltic Columns at the Giant's Causeway," by Profs. Jellett and O'Reilly.

MEETINGS FOR THE WEEK.

MONDAY, 9th.—Royal Geographical, 8.30.

TUESDAY, 10th.—Royal Institution, 3. "The Intellectual Movement of Germany from the Middle of the Last to the Middle of the Present Century," Prof. Hillebrand.

— Photographic, 8.
— Anthropological, 8.

WEDNESDAY, 11th.—Geological, 8.
— Microscopical, 8.

THURSDAY, 12th.—Royal Institution, 3. "The Intellectual Movement of Germany from the Middle of the Last to the Middle of the Present Century," Prof. Hillebrand.

— Royal Society Club, 6.30.
FRIDAY, 13th.—Royal Institution, 9. "The Thunderer Gun Explosion," Mr. F. J. Bramwell.

SATURDAY, 14th.—Physical, 3. "On the Suppression of the Induction Disturbance of the Telephone," Prof. H. McLeod; "On the Sensitive State of Electric Discharges through Rarefied Media," W. Spottiswoode and J. F. Moulton; "On a New Measuring Polariscopes," Prof. W. G. Adams.

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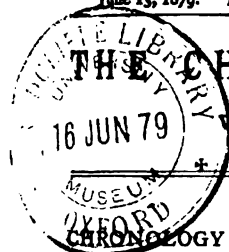
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THE CHEMICAL NEWS.

VOL. XXXIX. No. 1020.

NOTES ON THE CHRONOLOGY OF THE ISOMERIC PURPURINS, AND THE ACTUAL RELATIONS OF THE BODIES WHICH HAVE BEEN CALLED BY THE NAMES ANTHRA-PURPURIN, ISO-PURPURIN, AND FLAVO-PURPURIN.

By HENRY MORTON, Ph.D.,
President of the Stevens Institute of Technology.

THERE is a piece of recent chemical history which has already become much involved on account of the oversight of some writers, and as the threads happen to be in my hands, I think I shall do a good office to future students by laying these out straight at the present time.

In 1870 Mr. W. H. Perkin published in the *Journal of the London Chemical Society* a paper entitled "On Artificial Alizarin" (vol. viii., p. 143), in which at the end, in a footnote, the author says—"When purifying artificial alizarin by converting it into an alumina lake, I found that upon digestion with carbonate of potash this lake gave a red-coloured solution containing a colouring matter, dyeing mordants very similarly to alizarin, with this difference, that the reds are more scarlet and the purples bluer or more slaty. I have not obtained this body in a perfectly pure state as yet, but it appears to be crystalline."

The investigation thus opened by Perkin was diligently prosecuted by him, and at the meeting of the London Chemical Society, held June 6, 1872, he read a paper on this body, an abstract of which paper is published in the *CHEMICAL NEWS* (vol. xxv., p. 284), and a fuller report of which appears in the *Journal of the London Chemical Society*, vol. xxv., p. 659, under the title "Note on a Secondary Colouring Matter Produced in the Preparation of Alizarin from Anthracen." In these publications he gives the formula of the new body as $C_{14}H_6O_5$, and notes its relations and differences to and from the purpurin of madder. In the next volume of the *Proceedings*, or rather *Journal of the Chemical Society*, viz., vol. xxvi., 1873, Perkin published on page 425, under the title "On Anthrapurpurin," a long article on the same substance, to which were attached, as illustrations, pieces of cloth dyed with the substance as well as others dyed with alizarin. This article has been translated and quoted extensively in other publications, but the other articles seem to have been overlooked or forgotten. Hence have arisen some of the errors which I now propose to correct.

From the above references it will be seen that Perkin first separated out and recognised this new body in 1870, and diligently following up the research, published its formula in the beginning of June, 1872, and a full and exhaustive memoir in 1873.

In the *Moniteur Scientifique* for August, 1872, or just two years after the first publication by Perkin, and just two months after his second, in which he gave the formula of his new substance, appeared a communication announcing the discovery of a new product extracted from artificial alizarin by G. Auerbach, and named by him isopurpurin.

No mention whatever was made in this article of Perkin's prior publication, and it was of course generally inferred that Auerbach knew of them, but regarded his isopurpurin as a new discovery of his own, and not a mere confirmation of Perkin's work. In the course of time it appeared, however, that this was not the case, but that Auerbach

was ignorant of Perkin's prior publications, and, believing his material to be the same as that more fully described by Perkin in 1873, claimed priority.

Into this error he seems to have led several writers. Thus, Graebe and Liebermann, in their report on Artificial Alizarin at the Vienna Exhibition, under the heading isopurpurin, refer first to Auerbach's publication in the *Moniteur Scientifique* of August, 1872, and then to Perkin's third publication in the *Journal of the Chemical Society* of 1873, omitting all reference to, and evidently not knowing of, his earlier publications in 1870 and June of 1872.

Rosenstiehl, in a recent article, has fallen into a like error, and credits Auerbach with first obtaining this body in an impure state, and Perkin with having first extracted it pure in 1873.

But admitting that Auerbach is right in assuming that what he extracted was only the anthrapurpurin of Perkin, it is evident that he has no claim whatever to priority, but that the substance must be credited to Perkin, and should be called by his name of Anthrapurpurin.

As a matter of fact, however, the substance actually extracted by the method described by Auerbach in 1872 is not anthrapurpurin, but a mixture consisting largely of a substance which was new at that time, but was afterwards isolated by Messrs. Schunck and Roemer.

My reasons for this opinion are the following:—In the latter part of 1874 I had occasion, together with my friend Mr. Wm. E. Geyer, B.S., to repeat both Perkin's and Auerbach's method of treatment upon considerable quantities of artificial alizarin. We first followed out Perkin's method, and obtained a product corresponding in all respects with his description of anthrapurpurin. We then worked on another quantity of artificial alizarin according to Auerbach's directions, and when we came to the last treatment, namely, solution in and crystallisation from alcohol, we found that we evidently had to do with a mixture of two bodies differing in a marked way as to their solubilities.

Bearing in mind that we had found Perkin's anthrapurpurin but little soluble in alcohol, and that Auerbach described his isopurpurin as quite soluble, we so carried on the alcohol treatment as to reserve the soluble part as isopurpurin and reject the less soluble portion.

When we had carried this separation to a sufficient extent, we found that we had a body entirely distinct from Perkin's anthrapurpurin, but sufficiently like the description given by Auerbach of his isopurpurin to answer for it when due allowance was made for the uncertainties of a verbal description. When, however, the paper of Schunck and Roemer appeared in 1875, we saw that the body we had extracted agreed still more closely with their flavopurpurin.

Having obtained it, however, by the process described in 1872 by Auerbach, I still continued to call it by his name, and should do so still had he not very emphatically repudiated all credit for and connection with it.

The true relation of the substances which have borne the names given at the head of this article is, then, this:—

Anthrapurpurin is a single substance, first isolated by Perkin, partially in 1870, and thoroughly prior to June 6, 1872.

Isopurpurin is a mixture, in proportions varying according to the composition of the artificial alizarin from which it is extracted, and the way in which the alcoholic treatment is conducted, of anthra- and flavopurpurin with a trace of alizarin. This mixture was first separated and described by G. Auerbach in August, 1872.

Flavopurpurin is a single substance or chemical individual first isolated and described by Schunck and Roemer in the *Ber. der Deut. Chem. Gesell.*, 1876, p. 678.

The facts as to the composition of the material obtained by treating artificial alizarin by the method described by Auerbach are also established by the recent researches of M. A. Rosenstiehl, presented to the Chemical Society of Paris, and published in the *Bulletin* of that body of May 5 and 20, 1878, where, in vol. 29, p. 408, I find as follows:—

"On submitting isopurpurin obtained by the method described by M. Auerbach to proximate analysis, I was not slow in discovering the presence of 15 per cent of alizarin." Here follows a long and minute description of how this was removed by treatment with hot benzol.

The material so purified was then dissolved in boiling alcohol, and allowed to deposit crystals on cooling. The author then continues:—

"Comparative dyeing tests with the part crystallising out and with that remaining in the mother-liquor showed that the alcohol eliminated a substance colouring mordants like flavopurpurin, that is to say, of a red more orange than that of oxanthraflavon."

It should be mentioned that in this essay its author indicates by the word oxanthraflavon the material isolated by Perkin's method, but which through oversight of the earlier publications he credits to Auerbach, and names isopurpurin in consequence of this mistake.

ON THE COMPOUNDS OF THE TERPENES WITH HYDROCHLORIC ACID.

By WILLIAM A. TILDEN.

IN the course of researches which I have for some time been pursuing into the constitution of the terpenes I have had occasion to prepare and examine the compounds which these hydrocarbons form with hydrochloric acid. The following is a brief account of the chief results arrived at in this direction, and as they possess some interest at the present time, I think it desirable to place them on record.

When dry hydrochloric acid gas is passed into cooled turpentine oil (b.p. 156°), whether from American or French turpentine, the well-known mono-hydrochloride, $C_{10}H_{17}Cl$, is obtained, melting at 125°, and boiling at about 210°. It is accompanied by liquid products, which yield a large quantity of the same hydrochloride on distillation. This liquid consists of a mixture of cymene, always present in turpentine oil, with liquid compounds of the mono-hydrochloride and di-hydrochloride. The fact that the two chlorides when mixed together unite and liquefy was originally pointed out by Berthelot, and I can confirm this observation. The existence of a liquid modification of the mono-hydrochloride is doubtful. The solid mono-hydrochloride is also produced if the turpentine is diluted with benzol or carbon bisulphide before submitting it to the action of the gas.

The result is different when the turpentine is saturated with hydrochloric acid in the presence of some liquid which contains water, or is capable of yielding the elements of water. Thus, when the same turpentine oil is diluted with alcohol, ether, or glacial acetic acid, it yields a deep purplish coloured liquid, which, after exposure to the air for some time, deposits crystals of the di-hydrochloride, $C_{10}H_{18}Cl_2$. But it deserves to be noticed that when once the terpene has been made to unite with one molecule of hydrochloric acid no further action of the gas, either in the dry state or in presence of any of the solvents named, is capable of causing it to combine with a second molecule so as to produce the di-hydrochloride. There can be no doubt that these two chlorides are of entirely dissimilar constitution, as they behave quite differently under the influence of alkalies, the mono-hydrochloride being decomposed with difficulty by soda, yielding a crystalline camphene, whilst the di-hydrochloride produces chiefly terpinol, $C_{10}H_{17}(OH)$. The di-hydrochloride referred to possesses the same properties, whether prepared from lævo- or dextro-rotatory turpentine oil. Its solution in alcohol is optically inactive. The crystals melt at 48° in an atmosphere of hydrochloric acid, and by heat are readily decomposed into hydrochloric acid and a hydrocarbon. This di-hydrochloride is also obtained by the

action of hydrochloric acid gas upon crystallised terpin hydrate and upon terpinol. (*Journ. Chem. Soc.*, June, 1878.)

If in place of turpentine oil one of the terpenes of higher boiling-point is operated upon no mono-hydrochloride can be obtained. Thus, the terpene from orange oil boiling at 176° was dried and saturated with hydrochloric acid. The resulting solution was colourless, and after exposure to the air for a day or two was found to contain no chlorine at all. But when the same terpene was mixed with ether, and treated with the gas, a solid mass of colourless crystals remained after evaporation of the ether. It was a somewhat unexpected discovery that these crystals consist of the same di-hydrochloride already described, that is obtained under the same circumstances from turpentine. There is, however, some difference observable in the behaviour of the terpene from the orange and those from the turpentine oils; for whilst the former yields a nearly theoretical amount of crystallised di-hydrochloride, the latter produce a considerable quantity of liquid of deep colour. This colouration is characteristic of the action of hydrochloric acid upon terpinol, $C_{10}H_{18}O$, or $C_{10}H_{17}OH$, as shown in a recent paper (*Journal of the Chemical Society*, June, 1879), and it appears to indicate the formation of an intermediate product which is probably the same in both cases.

In order further to establish the identity of the di-hydrochlorides thus obtained from the two classes of terpenes, the dichloride prepared from the orange oil and from turpentine oil were decomposed by water. 50 grms. of each boiled with 500 c.c. of water were found to be decomposed at the same rate and in the same manner, yielding a mixture of hydrocarbon (terpinylene, $C_{10}H_{16}$, b.p. 176°), and terpinol ($C_{10}H_{18}O$, b.p. 210°). The terpinol was recognised in each case by its conversion into crystallised terpin by the action of sulphuric acid and water. The terpenes from juniper oil (b.p. 156°), lemon (b.p. 176°), bergamot (b.p. 176°), and caraway (b.p. 176°) yield this di-hydrochloride, as also does the terpene (b.p. 174° to 176°) contained in the *ol. pini sylvestris* of pharmacy and a terpene (b.p. about 176°), obtained from rosin spirit.

The single exception is the "sylvestrene" discovered by Atterberg in Swedish turpentine, and which I have since recognised in Russian turpentine. The di-hydrochloride which this terpene yields melts at 72° to 73°, and although decomposed by alkali it yields a hydrate which is apparently not common terpinol. If it should prove to be distinct from ordinary terpinol it is probable that it may be convertible into a new terpin. I am about to make experiments with the object of settling this question, which possesses some interest.

The terpene di-hydrochloride described in this paper is decomposed by simple application of heat, and by prolonged boiling in a flask fitted with condensing tube is resolved into hydrochloric acid and a hydrocarbon, terpinylene, $C_{10}H_{16}$, which boils at 176°. A portion of the hydrocarbon is at the same time converted into viscid polymers. Terpinylene cannot, however, be made to recombine with hydrochloric acid so as to reproduce the crystalline di-hydrochloride.

Thus terpenes of the two classes, boiling respectively at 156° and 176°, and very diverse in their optical properties, may all, by this process, be converted into one and the same inactive hydrocarbon.

I cannot conclude without expressing my thanks to Mr. G. H. Morris and Dr. G. Harrow for valuable assistance in these experiments and in other parts of the same research.

The Paris Academy of Sciences.—At the sitting of the Academy on Monday last Professor G. G. Stokes, M.A., D.C.L., LL.D., Secretary of the Royal Society, was elected a corresponding member in the Physical Section in succession to the late M. Angström.

ON NITROGEN IODIDE.

By J. W. MALLET.

THE various researches upon the highly explosive black substance formed by the action of iodine upon ammonia have, as is well known, led to different views of its composition. That first put forward by Colin and Gay-Lussac, namely that it is nitrogen tri-iodide, appeared to have been set aside by the experiments of Serullas,* Millon,† Bineau,‡ Gladstone,§ Playfair, and Bunsen,|| all of whom found hydrogen to be a constituent, but in varying amount, the formulæ $\text{NH}_3\text{I}-\text{NH}_2\text{I}-\text{NH}_3\text{I}$ and $\text{NH}_3\cdot 4\text{NI}_3$ having been assigned to the products obtained by somewhat different processes, until more recently Stahlschmidt¶ has revived the statement that under certain conditions at least the simple tri-iodide is formed.

One source of error in the analysis of this instable compound seems to have received insufficient attention from some of those who have examined it, and somewhat affects the conclusions drawn from the experiments of others. The substance, obtained as a black powder by the action of iodine on ammonia, decomposes gradually in contact with water, nitrogen gas being slowly evolved, iodine liberated, and iodic and hydriodic acids or ammonium salts formed. This decomposition takes place to no inconsiderable extent during the drying of the powder at ordinary temperature, and may be readily proved to have occurred by treating the dry residue with any of the solvents for iodine, as for instance alcohol, chloroform, or carbon di-sulphide, which at once form deeply coloured solutions. Hence the air-dried powder is really a mixture of the original substance or substances with various decomposition products, and so is unsuited for accurate analysis. On the other hand, it might be doubted in regard to the analyses made of freshly prepared and still moist iodide, whether the results apply to a substance capable of being obtained in the dry state without decomposition.

Millon and Bineau, it is true, seem to have aimed at guarding against this difficulty by drying the materials, side by side with solid potash, under a jar of gaseous ammonia. Bineau remarks that some of the gas is at first absorbed by the water moistening the powder, but that as the drying proceeds the gas recovers ("à peu près") its original volume, and that so moisture is gotten rid of without the iodide being decomposed. But, without qualitative examination of the gas, or a very accurate determination of its volume, this does not afford satisfactory proof that the dry substance is unchanged. I have observed continuous, though slow, evolution of little bubbles of nitrogen from the freshly prepared powder when kept under strong aqueous ammonia, though in this case no free iodine can be found, since as fast as it is liberated it reacts with the excess of ammonia, reproducing the nitrogen iodide, and forming more ammonium iodide, which latter therefore tends to accumulate in the liquid, and is thus to be found in the residue from drying up a moist mass of the powder in an atmosphere of ammonia.

I have applied a different method, which is often useful under other circumstances, for quickly getting rid of water, namely repeated and rapid washing with absolute alcohol, followed by ether, and final evaporation of the last-named volatile liquid.

20 or 30 grms. of iodine was dissolved in a minimum of 95 per cent. alcohol, and precipitated by pouring into a large volume of cold water. The finely divided iodine was washed several times by decantation, then gently triturated for several minutes in a porcelain mortar with

a large excess of the strongest liquid ammonia kept at or below 0°C . by a freezing mixture, the liquid poured off from the easily subsiding black powder, and replaced two or three times by fresh solution of ammonia. The powder was then at once transferred to a corked flask, and shaken up repeatedly, first with alcohol of 95 per cent., then with absolute alcohol, and finally with anhydrous ether, all of these liquids being artificially cooled. Most of the last portion of ether, which, as well as several of the preceding washings, was perfectly colourless, having been decanted off, the fluid black mud was turned out upon a filter, drained in a few moments, and the remains of the ether swept away as vapour by placing the filter and contents under a receiver and drawing cold dry air through in a rapid stream. This use of alcohol and ether to remove water from the iodide after its formation is not to be confounded with the employment by several experimenters of alcoholic solutions of iodine or ammonia, or both, in making the substance in the first instance.

The product thus obtained was explosive in the highest degree, yielding to very slight friction upon paper, frequently producing a crackling sound from little partial explosions when it was gently rubbed under water, and in two instances exploding in some quantity under water, with much violence and complete shattering of the vessel.

The relative quantities of nitrogen and iodine were ascertained by decomposing an unweighed parcel of the substance by a moderately strong aqueous solution of sodium sulphite, neutralised by dilute sodium hydrate, and determining, in equal volumes of the liquid, the nitrogen as ammonia (expelled by boiling with excess of soda, and collected in sulphuric acid of standard strength), and the iodine as silver iodide.

A—In three experiments were obtained—

	No. 1.	No. 2.	No. 3.
NH_3	0.126	0.184	0.172 grms.
Equiv. to N	0.104	0.152	0.142 "
AgI	5.126	7.607	6.889 "
Equiv. to I	2.771	4.112	3.723 "

Nos. 1 and 2 refer to the same lot of material (separately decomposed, however); No. 3 to the product from a repetition of the same process.

These numbers correspond to—

	No. 1.	No. 2.	No. 3.	Calc. for NI_3 .
N (assumed)	14.0	14.0	14.0	14.0
I	373.02	378.74	367.04	381.0

or, for 1 atom of nitrogen—

	No. 1.	No. 2.	No. 3.
Atoms of iodine ..	2.94	2.98	2.89

B—Another specimen, similarly prepared, but with weaker ammonia, without any precaution as to cooling the materials used, and in a room the temperature of which was about 23°C ., gave—

	No. 1.	No. 2.
NH_3	0.174	0.152 grms.
Equiv. to N	0.143	0.125 "
AgI	5.935	5.467 "
Equiv. to I	3.208	2.955 "

corresponding to—

	No. 1.	No. 2.
N (assumed)	14.00	14.00
I	314.07	330.96

or, for one atom of nitrogen—

	No. 1.	No. 2.
Atoms of iodine	2.47	2.61

C—Two other specimens, washed at first, not with alcohol and ether, but simply with water until no free ammonia could be detected in the washings, then kept under water for two or three days at ordinary temperature, with occasional stirring, and finally dried after washing out with absolute alcohol and ether, as above described, gave—

* Ann. Ch. Phys. [2], 42; 200 (1829).

† Ann. Ch. Phys. [2], 69; 78 (1838).

‡ Ann. Ch. Phys. [2], 70; 270 (1838); and [3], 15; 71 (1845).

§ Chem. Soc. Q. Journ., 4; 34 (1851).

¶ Ann. Ch. Pharm., 84; 1 (1852).

¶ Pogg. Ann., 119; 421 (1863).

	No. 1.	No. 2.
NH ₃	0.136	0.127 grms.
Equiv. to N ..	0.112	0.105 "
Ag I	3.954	3.667 "
Equiv. to I ..	2.137	1.982 "

corresponding to—

N (assumed) ..	14.00	14.00
I	267.12	264.27

or, for 1 atom of nitrogen—

Atoms of iodine ..	2.10	2.08
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Of the above results—

A distinctly represents the atomic ratio N:I=1:3, fully confirming Stahlschmidt's conclusion that the tri-iodide can be obtained quite free from hydrogen.

B corresponds well with N:I=2:5, and

C with the proportion found as the result of several earlier experiments, N:I=1:2.

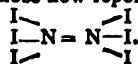
If B be admitted to stand for a definite compound, and not a mere mixture of A and C, which is rendered probable by the fairly close agreement of the results obtained with the calculated figures, there must be two atoms of nitrogen in the molecule, and the formula will be N₂HI₃. This formula may almost as well be deduced from Bunsen's analysis of one of the products he obtained (by addition of ammonia to a solution of iodine in nitro-hydrochloric acid, diluted with water, and rapidly washing with cold water) as that which he has himself assigned, viz., NH₃.4NI₃, since the figures stand—

	Calc. for N ₂ HI ₃ .	Calc. for NH ₃ .4NI ₃ .	Found by Bunsen.
N 4.22	4.38 ..	0.008143 or assumed 4.38	
H 0.15	0.19 ..		
I 95.63	95.43 ..	0.174417	93.82*
100.00	100.00		

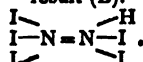
It will be seen that the figures of the formula now proposed differ less from those of Bunsen's formula than the latter do from the results of experiment.

In view of the general fact that the compounds of nitrogen in which this element behaves as a pentad are those in which instability is chiefly observable, and noticing the various proportions in which the iodine and hydrogen have been found together united with it, it seems fairly probable that the molecule of each of these explosive compounds contains two pentad nitrogen atoms; and reviewing all that has been published on the subject, with attention to the sources of error connected with the various methods used, we seem to have established the following series of substituted products, beginning with the tri-iodide:—

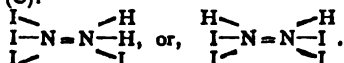
Experiments of Stahlschmidt (using aqueous ammonia) and those now reported (A).



Experiments of Bunsen (second product), and above result (B).

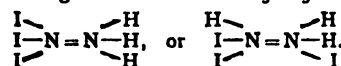


Experiments of Bineau, Stahlschmidt, Gladstone (using alcoholic solutions of ammonia and iodine), and above result (C).



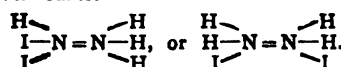
* There seems to be an error in the calculation of the percentage of iodine in the original paper (*Ann. d. Ch. u. Pharm.*; 84; 1), but the figures here given are correctly deduced, I believe, from the actual weighings quoted, viz., 0.1300 grm. of ammonio-chloride of platinum and 0.0732 grm. of palladium.

Experiments of Bunsen (first product, using anhydrous alcoholic solutions of ammonia and iodine). He himself assigned the formula NH₃.NI₃.



It is much more doubtful whether we should add—

Experiments of Milon (in 1838). The product dried under a receiver filled with gaseous ammonia had lost its extreme explosiveness, and no doubt contained ammonium iodide.



It appears that, not only may these different products be obtained by varying the conditions under which the formation of the explosive substance is effected, but during its decomposition by water, certainly involving more than a single series of reactions, one of these products may be more or less converted into others.—*American Chemical Journal*.

University of Virginia, Feb. 21, 1879.

ON THE SOFTENING OF MAGNESIA-HARD WATER.

By J. GROSSMANN, Ph.D.

I HAVE lately had an opportunity of making some experiments on the softening of so-called magnesia-hard water by Clark's process. Although from a theoretical point of view there is no reason why such water should not be softened as easily as ordinary (lime-)hard water, it was thought convenient to try the experiment on a somewhat large scale, especially as I have not been able to find that any experiments on the softening of magnesia-hard water have been published before in detail.

The water in question is pumped from a colliery at Collins Green, near St. Helens. The bottom of the well is about 96½ yards from the ground, and I give for completeness sake the different strata, as well as the quantity of lime and magnesia per hundred, which hydrochloric acid dissolved out of them, the samples being previously dried at 240° F.

	P.c. CaO.	P.c. MgO.	Yards.
Depth of alluvium and clay from surface of ground to top of the red sandstone	—	—	17
New red sandstone	0.16	0.24	34
Yellow sandstone	0.10	0.22	13
Compact red sandstone	4.72	0.37	12
New marl or metal	2.16	0.52	6
Dun Rock	0.40	0.46	1
Coarse porous red sandstone ..	1.88	0.27	13
Coarse red sand	0.72	0.32	1
			96½

This water is organically very pure and contains only small quantities of chlorides and sulphates. The hardness is almost entirely temporary, though in estimating the permanent hardness it is sometimes difficult to remove the lime and magnesia by boiling. I may as well point out in this place a common fallacy, which is that hard water by being boiled before being used for domestic purposes becomes soft. As it is generally only heated to boiling just before being used for washing or cooking it makes not much difference in the hardness of the water. The following is taken from the Sixth Report of the Commissioners appointed in 1868 to inquire into the best

means of preventing the pollution of rivers. Thirteen samples of water drawn on thirteen different days from the kitchen boiler of a dwelling-house, and from that of the Athenæum Club, were found to be usually nearly as hard as the cold water with which those boilers were supplied, as is seen from the following results:—

Hardness of Cold Water.	Hardness of Hot Water.	Hardness of Cold Water.	Hardness of Hot Water.
14.6	13.6	15.4	14.3
14.4	13.9	15.9	11.9
14.4	13.4	16.1	11.9
14.6	11.6	18.7	18.4
14.6	7.6	18.7	18.6
14.4	11.7	18.7	18.4
14.4	12.1	—	—

I boiled Collins Green water that tested 27.8° total hardness in an ordinary tea-kettle for five minutes, and even that exceptionally temporary-hard water tested then still 17.6° total hardness. The experiments for softening this water by Clark's process were made in two series; in the first milk of lime was used, and in the second lime-water. The milk of lime was prepared by mixing 5 lbs. of slaked lime of 70 per cent CaO with 6 gallons of Collins Green water. In performing the softening of the water the milk of lime was gradually added to the water during the operation, the mixture being well agitated during the whole experiment.

In three experiments the hardness was reduced from 23.3° which the water tested originally to 5.0°, 5.5°, and 5.7°. The total solids were reduced respectively from 33.9 to 13.0, 33.6 to 12.2, and 33.6 to 13.6 grs. per gallon. The softened water, after it had been standing in stoppered bottles for a few days, formed a little deposit, and its hardness was then 3.5° in all cases. The analysis showed that then the lime had been reduced from 9.13 to 1.62; the magnesia from 5.23 to 0.23 grs. per gallon.

There were used on an average 1465 gallons of water to 3.5 lbs. of CaO, corresponding to 2.67 lbs. of lime of 90 per cent per 1000 gallons. Theoretically to precipitate the lime and magnesia as far as I have done in my experiments would require 2.31 lbs. of CaO per 1000 gallons.

The experiments for softening with lime-water were carried out in a similar way. A measured quantity of the lime-water was put into the mixing tank, and the Collins Green water run into it until nearly all the lime of the former was taken up, which in these experiments as well as in those with milk of lime was ascertained by the well known silver test. The mixture was well agitated during the whole experiment. A great advantage in working with lime-water is that it requires less agitating than working with milk of lime.

In four experiments the hardness was reduced from 23.3°, which the water tested originally, to 3.5°, 3.3°, 3.5°, and 3.5°. The total solids were reduced respectively from 34.3 to 12.0, 34.4 to 11.2, 34.6 to 12.0, and 34.7 to 11.2. The lime had been reduced from 9.07 to 1.67, and the magnesia from 5.24 to 0.18 grs. per gallon.

There were used on an average 1670 gallons of water on 530 gallons of lime-water, which latter having been exposed to the air for some time before being used tested only 60 grs. CaO per gallon. This corresponds to one volume of lime-water of 60 grs. CaO per gallon to 3.13 volumes of Collins Green water, or corrected to 90 grs. of CaO per gallon, which strong lime-water should contain, the proportion of lime-water to Collins Green water would be 1 to 4.72. This result is not quite correct, as the lime-water had undergone a further susceptible decomposition while lying in the mixing-tank; so that at the time of its action on the Collins Green water it contained less than 60 grs. CaO per gallon. Both in working with milk of lime and lime-water the softened water settled perfectly clear after three to five hours, taken from the time that the agitation was suspended.

From these experiments it follows that magnesia-hard

water softens as well and as easily by Clark's process as ordinary (lime-)hard water.

I wish to point out here that in my opinion, with our present knowledge of the behaviour of magnesia-hard water towards soap solution, the hardness test in cases of water of that description cannot be relied on in any way. Mr. Wanklyn, in his book "On Water Analysis," says that magnesia takes up as much soap as 1½ equivalents of lime would take up. If that were so the Collins Green water should have tested 35.9° total hardness, instead of 23.3°, which it gave on the most careful testing. It will be noticed that this is even less than the water would require if it contained an equivalent of lime for the magnesia present; for the calculated hardness would then be 29.4°. It appears to me that in the case of magnesia-hard water a determination of the lime and magnesia is the only means of showing the quality of the water. On the other hand, I quite agree with Mr. Wanklyn in putting little reliance on the distinction between temporary and permanent hardness, unless the estimations have been performed very carefully; and there are cases in which, even with the most careful working, figures for temporary and permanent hardness may be got which are not trustworthy.

ANALYSIS OF THE WATER OF ST. DUNSTAN'S WELL, MELROSE.

By WILLIAM JOHNSTONE, F.C.S., F.I.C.

An exhaustive analysis of this strongly ferruginous saline carbonated water has not, so far as I have been able to learn, been published before, although a general analysis of it was made by Mr. J. Dewar in 1870, and published in the CHEMICAL NEWS, vol. xxiv., p. 171. There has been a partial analysis of this water also by Dr. Stevenson Macadam, of Edinburgh, but as his results are very doubtful I refrain from quoting it at all. As this water is peculiar for the large quantity of iron it contains, and as its composition seems to have varied considerably since it was examined by Prof. Dewar, it is thought that another and more complete analysis will not be out of place.

Melrose is a village of some 2000 inhabitants, and is situated at the base of the Eildon Hills, in the valley of the Tweed; one of those lovely spots that are said to be particularly favourable to the curative effects of medicinal waters. The surrounding country consists principally of richly variegated fields, and the entire district teems with objects of natural beauty, while every spot on which the eye rests is associated with some incident of historic or classic interest—the ecclesiasticism and chivalry of the past, and the interest conferred by Scott in the present. The salubrity of the district is well known, and the romantic scenery, which is equalled by few places in the kingdom, presents a picture dear to the lover of nature as well as to the invalid.

The spring, which belongs to Mr. John Turnbull, of St. Dunstan's Cottage, Melrose, was accidentally discovered in 1870, when sinking a shaft for ordinary spring water. The water is said to be uniform in quantity, its temperature varies only within very narrow limits, and is quite cold. As it occurs in the spring it is perfectly clear, of a yellowish colour, but, after exposure to the air, deposits the iron as a ferric manganic oxide; it has a strong powdery odour and a decided inky taste. When taken from the well there is a disengagement of small pearly bubbles, and on being shaken up in a closed bottle liberates a large quantity of gas. The temperature of the spring on October 5, 1878, was 50° F., the temperature of the air at the same time being 62° F. The specific gravity of the water is 1.0019356. A careful qualitative analysis indicated the presence of the following constituents in estimable quantities:—

Bases.	Acids, or Elements replacing them.
Soda	Chlorine
Potash	Sulphuric acid
Lithia	Phosphoric "
Ammonia	Carbonic "
Lime	Silicic "
Magnesia	Crenic "
Baryta	Apocrenic "
Strontia	Bromine "
Ferrous oxide	Iodine
Manganous oxide	
Alumina	

In addition to the above were found traces of fluorine, butyric and phosphoric acids, along with other volatile and non-volatile organic matter.

The residue of 30 gallons was examined by means of the spectroscope for caesium, rubidium, and thallium, but the search proved unsuccessful. The method of quantitative analysis employed was essentially that described by Fresenius under the head of "Analysis of Mineral Waters" in the last edition of his "Manual of Quantitative Analysis."

The following figures represent grammes per litre, and are the average of three determinations of each of the principal constituents, done in order to guard against error and to ensure accuracy; the amount of water taken for the several determinations was, whenever practicable, ascertained by weight.

Results of Analysis.

Chlorine	0'378243
Bromine	0'000305
Iodine	0'000788
Sulphuric acid (SO ₃)	0'074258
Carbonic dioxide (total)	0'682183
Silicic acid	0'035286
Ferrous oxide.. .. .	0'328680
Lime and strontia expressed as carbonates	0'103842
Magnesia (total)	0'056216
Lime precipitated as carbonate on boiling.. .. .	0'028544
Baryta	0'002917
Strontia	0'000886
Manganous oxide	0'000785
Phosphoric acid	0'020085
Alumina	0'006582
Lithia	0'000282
Potash	0'006258
Soda.. .. .	0'343063
Ammonia	0'018652

Total fixed constituents 1'390000

Following the arrangement adopted by Fresenius on the assumption that the strongest acids are united with the strongest bases, &c., and allowing for the fact that the more or less degree of the solubility of the salts decidedly influence the manifestation of the affinities, and in order to simplify reference and comparison, and by way of contrast, I append the results calculated as grammes per litre. (See next column.)

Gases dissolved in the water and expelled by ebullition *in vacuo* measured at 59° F. and 760 m.m. barometer.

	Cubic centims.
Carbonic dioxide	101'434
Oxygen	6'793
Nitrogen	31'250

139'477

I am unable at present to form an opinion as to whether the composition of the water is constant, owing to the surface-water having had free admittance to the well previous to the collection of this sample on March 18, 1879,

Barium sulphate	0'004441
Strontium sulphate	0'001571
Calcium sulphate	0'088017
Magnesium sulphate	0'030377
Magnesium bromide	0'000397
Magnesium iodide.. .. .	0'000945
Potassium chloride	0'009900
Lithium chloride	0'000463
Ammonium chloride	0'040985
Sodium chloride	0'544276
Magnesium chloride	0'027528
Aluminium phosphate	0'019503
Calcium phosphate	0'012511
Calcium carbonate	0'028544
Magnesium carbonate	0'037205
Ferrous carbonate.. .. .	0'529553
Manganous carbonate	0'001278
Silicic acid	0'027866
Crenic acid	0'013572
Apocrenic acid	0'000487

Total ingredients calculated 1'420519

but at the same time I may mention that a considerable sum of money has been expended in the view of preventing any more surface-water entering the well, so hope at some future date to be able to favour you with another analysis.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, June 5, 1879.

Mr. WARREN DE LA RUE, President, in the Chair.

AFTER the transaction of the usual business the following certificates were read for the first time:—E. Buckney, R. E. Holloway, T. Blackburn, E. F. Mondy, E. Francis. It was announced that a ballot for the election of Fellows would take place at the next meeting of the Society (June 19).

The following papers were read:—

"A Contribution to the Theory of Fractional Distillation," by T. E. THORPE. Wanklyn some years ago found that when two liquids of different boiling-points were mixed together in equal quantities by weight and distilled, the proportion of each constituent in the distillate was the product of its vapour-density and vapour-tension at the temperature of ebullition of the fraction. Hence, under certain circumstances the less volatile of the two substances may pass over most rapidly, whilst if the vapour-tensions and vapour-densities of the two liquids are inversely proportional the mixture will distil unchanged. Berthelot observed that a mixture of 90'9 parts of carbon disulphide with 9'1 parts of ethyl alcohol boiled and distilled as a homogeneous liquid. An instance of this phenomenon has been noticed by the author. A mixture of equal volumes of carbon tetrachloride boiling at 76'6°, and of methyl alcohol boiling at 65'2°, was distilled. It was found that 46'5 per cent of the whole boiled constantly between 55'6° and 55'9°, 10° lower than the boiling-point of the most volatile constituent. This mixture contained 78'1 per cent carbon tetrachloride and 21'9 per cent methyl alcohol. This proportion, 3'6 to 1, is almost identical with that obtained by multiplying the vapour-tensions of the two liquids at the temperature of the boiling-point of the mixture (55'7°) by their respective vapour densities, $487'4 \times 15'97 : 372'4 \times 76'69 :: 1 : 3'67$. The distillation of the residue in the flask was continued by Mr. C. C. Starling: at first principally carbon tetrachloride, finally pure methyl alcohol passed over. The author suggests as a striking lecture experiment the following:—Three baro-

meter tubes are filled with mercury. Into one some methyl alcohol is passed; into the second some carbon tetrachloride; into the third a mixture containing by volume 3 parts of methyl alcohol to 5 of carbon tetrachloride: the depressions of the mercury column are 80, 70, and 130 m.m. respectively. The author promises a further research on the physical peculiarities of this mixture.

"Preliminary Note on the Action of Organo-zinc Compounds on Quinons," by F. R. JAPP. When finely-powdered phenanthren quinon is gradually added to zinc ethyl, diluted with ether so as not to be spontaneously inflammable, a reaction takes place with evolution of gas. The orange colour of the quinon disappears: a whitish powder is formed, which sinks to the bottom of the liquid. On decomposing this product with an excess of alcohol, boiling, and filtering hot, transparent, faintly yellowish, rectangular plates were obtained, fusing at 77° , and having the formula $C_{18}H_{20}O_3$. This formula can be resolved into—



The compound $C_{16}H_{14}O_2$ has not yet been obtained pure; but the monacetyl derivative, $C_{16}H_{13}O_2(C_2H_3O)$, has been prepared and analysed; it fuses at 103° . The author refrains at present from discussing the constitution of the compound $C_{16}H_{14}O_2$, but suggests that these reactions may serve to distinguish quinons from double ketons. He intends, also, to study the action of organo-zinc compounds on other quinons and allied substances as dibenzoyl.

After some remarks by Dr. ARMSTRONG on the interest and probable bearing of the above reaction,

Dr. WRIGHT read a paper entitled "Third Report to the Chemical Society on Researches on some points in Chemical Dynamics (On the Curved Surfaces expressing the Relations between Time, Temperature, and Amount of De-oxidation of Copper Oxide by Hydrogen and Carbon Oxide)," by C. R. A. WRIGHT, A. P. LUFF, and E. H. RENNIE. This is a continuation of the previous reports by the authors on the subject. In the present paper a large number of observations have been made by reducing a uniform weight (1.15 grms.) of copper oxide (prepared by igniting pure copper nitrate) in narrow glass U-tubes heated to known and constant temperatures in vapour-baths: in some cases water and paraffin baths were employed fitted with Page's gas-regulator. Equable streams of hydrogen and carbonic oxide were obtained by bubbling the gas through a wash-bottle, the stream being adjusted by a screw-clamp, and counting the bubbles. The average rate was 12.5 c.c. per minute. By plotting out the results thus obtained in space with reference to three planes mutually at right angles, so that the distance from each plane represents, respectively, the time of exposure, the temperature, and the percentage loss of oxygen. Points are marked on curved surfaces, the sections of which, parallel to the three primary planes, represent respectively, the amounts of oxidation produced in given times at a constant temperature, the times required to produce given amounts of deoxidation at constant temperatures, and the amounts of deoxidation produced at given temperatures in a constant period of time. The paper was illustrated by tables, diagrams, and models of the curves thus produced, the mode of experimentation adopted being to determine for a constant temperature the amounts of deoxidation produced in varying times. At and above 160° in the case of hydrogen, and 130° in the case of carbonic oxide, the curious result was arrived at that the same mean curve is obtained whether the exposure be made all at once to a temperature for a time, T, or in periods of time t_1, t_2, t_3 , &c., which are together equal to T, provided that the interval between the periods is not too long (ten minutes). Below the above temperatures the deoxidation in an hour is greater than that produced in two consecutive heatings of half an hour each with an interval between. The curves obtained with hydrogen and carbonic oxide resemble each other in certain respects; their general cha-

rafter is that of a sigmoid. For a certain time there is no perceptible action; this time is the longer the lower the temperature. Reduction then commences languidly, quickly accelerating until a maximum of activity is reached, after which it diminishes until almost perfect deoxidation is effected. The maximum rate of action with hydrogen lies about 10 per cent (out of 19.74 per cent) of oxygen originally contained in the copper oxide, and about 7 to 8 per cent in the case of carbonic oxide. This apparently indicates that hydrogen passes through the outer and partially reduced surface of particles to their interior more readily than carbonic oxide. The following numbers illustrate the maximum rates of reduction attained, the gaseous currents being competent to remove 0.7791 per cent of oxygen (out of 19.74) per minute:—

Temperature.	Maximum rate with H. Per cent per minute.	Maximum rate with CO. Per cent per minute.
100°	—	0.135
118°	—	0.380
130°	—	0.480
160°	0.118	0.570
175°	0.220	—
184°	0.270	0.270
210°	0.490	—
256°	0.700	—

The higher the temperature the nearer does each curve approximate towards a limiting straight line, which would be attained did deoxidation commence immediately and go on at such a rate that all the hydrogen was converted into water, and the carbonic oxide into carbonic acid. The highest rates of reduction attained corresponded to a conversion of about nine-tenths of the H into H_2O , &c., of CO into CO_2 . From the curves it is evident that, *ceteris paribus*, to perform a given amount of deoxidation with hydrogen requires either a higher temperature or a longer time than with carbonic oxide. The existence of "a period of incubation" during which no action takes place, and the acceleration in the rate of action to a maximum, &c., shows that what has been termed "Chemical Induction" by Bunsen and Roscoe takes place in these cases to a large extent, dependent in amount on the temperature. From experiments now in progress this does not appear to be the case when copper is oxidised by hot air. A number of observations were made on the effect of varying the speed of the current of reducing gas and the weight of copper oxide used, with the general result that a more rapid stream or a smaller weight of copper oxide corresponds to an increased percentage of deoxidation, and *vice versa*. Heating the copper oxide just before use causes great irregularities in the action. A large number of observations were made by enclosing the copper oxide in sealed tubes filled with the respective gases, heating for different periods, and determining whether reduction had taken place or not; thus the following numbers were obtained. The times are perceptibly longer than those found in the corresponding U-tube experiments; in every case the time at any given temperature is less with CO than with H.

Temperature.	With CO, period of Incubation equals about—	With H, period of Incubation equals about—
160°	12 minutes	80 minutes
130	35 "	6 hours
118	$6\frac{1}{2}$ hours	12 "
100	7 $\frac{1}{2}$ "	28 "
83.84	11 "	180 "

Attempts were made to trace out curves illustrating the rate of reduction at various temperatures in atmospheres containing more reducing gas than would suffice to de-oxidise completely the CuO . The results were not very concordant, but indicated that the action here is still of the same kind; i.e., a period of incubation exists, after which action commences, reaches a maximum, and then falls off; in other words, chemical induction takes place whether the metallic oxide or the reducing gas be at any given momen-

in excess. In these experiments, too, either the time requisite to produce a given amount of action is less or the temperature is lower with carbonic oxide than with hydrogen. The paper, of which the above is but a brief extract, contains about 70 pages.

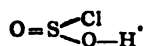
"On Fractional Distillation," by F. D. BROWN. The author considers the theory and formulæ involved in the process of distillation. He sums up as follows:—"The equation which represents the relation between the composition of the liquid and that of the vapour given off by it at a given pressure, together with that which represents the relation between the composition of the liquid and its boiling-point at the same pressure, contain all the experimental data which can be derived from the distillation, and form, together with the above formulæ, a complete history of it. If we can refer these relations to known laws we shall have arrived at an explanation of fractional distillation." The consideration of the distillation of mixtures divides itself into four heads—substances which are not miscible; substances which mix in all proportions but do not combine; substances which are soluble to a limited extent and do not combine; and substances which are mutually soluble and combine. The author then considers the researches of Magnus, Regnault, Pierre and Puchot, and A. Naumann as regards substances which do not mix. From these researches it was proved that the ratio of the molecules of the two liquids in the distillate is constant, and equal to that of their vapour-tensions at the temperature of distillation. The author has taken up the study of substances which mix in all proportions but which do not combine. The substances chosen were benzene and carbon disulphide. The composition of the mixture was ascertained by taking its density. These liquids expand on mixing; the greatest expansion occurs when the molecules of each are present in about equal numbers. In a series of thirteen tables the author gives the results of the fractional distillation of mixtures containing 61.95 per cent, 61.76 per cent, 70.86 per cent, 40.81 per cent, and 18.33 per cent of carbon disulphide respectively. The temperatures at which the fractions distilled, their weights and composition, and the composition of the liquid remaining in the still are given. The author exhibited some of the apparatus used in his researches.

In the discussion which followed, in which the President, Dr. Wright, Dr. Armstrong, and Mr. Friswell joined, there seemed some slight uncertainty as to the exact conclusions to be deduced from the author's experiments.

Mr. HOWARD pointed out that substances might dissolve in the liquid condition but not when in a state of vapour. Amylic alcohol would distil at 96° in the vapour of water, but the presence of a small quantity of ethylic alcohol would completely alter the composition of the vapour.

The two following papers were then read by the SECRETARY:—

"On Chlorstannic Acid," by J. W. MALLETT. A bottle containing a strong aqueous solution of stannous chloride, after standing for a year or two, deposited a transparent jelly-like substance of yellowish colour. This was washed and dried on a glass plate at the temperature of the atmosphere: it shrank up, cracked, and dried in fragments resembling gum arabic. Heated in a glass tube it evolved hydrochloric acid, leaving a white residue of stannic oxide free from chlorine. Its composition was SnO_2HCl , its constitution—



It formed salts with soda and ammonia. The author has not been able to reproduce this substance.

"On Indigo-purpurin and Indirubin," by E. SCHUNCK. Baeyer and Emmerling (*Ber. Deut. Chem. Gesell.*, iii., 514) described some years ago the formation of a red colouring matter with indigo-blue by the action of acetyl chloride, phosphorus trichloride, and phosphorus on isatin. This they named indigo-purpurin. Recently, Baeyer (*Ber. Deut. Chem. Gesell.*, xii., 457) describes another method of pre-

paring this colouring matter, and gives its properties. In 1856 the author read a paper (*Manchester Memoirs*, 2nd series, xiv., 181 to 237) and gave an account of a red colouring matter formed by the action of acids on indican, indigo-blue being formed. This colouring matter he named indirubin. Indigo-purpurin has all the properties of indirubin, and is, in fact, identical with it. The author therefore considers that the name indigo-purpurin should be abolished, and the original name indirubin retained.

The Society then adjourned to June 19, when a ballot for the election of Fellows will be taken, and the following papers read:—"On Gardenin," by Dr. Stenhouse and Mr. Groves; "On the Action of Sulphuric Acid on the Hydrocarbons of the Formula $\text{C}_{10}\text{H}_{16}$," by Drs. Armstrong and Tilden; "Researches on the Terpenes, Camphor, and Allied Compounds (Parts I. and II.)," by Dr. Armstrong; "Contributions to the History of Starch and its Transformations," by Horace T. Brown; "On the Boiling-points of certain Metals and Metallic Salts," by Dr. Carnelly and H. Carleton Williams; "On the Determination of Nitric Acid by means of Indigo," by R. Warington; "On Dry Copper-Zinc Couples and Analogous Agents," by Dr. Gladstone and Mr. Tribe; "Notes on the Purple of the Ancients," by R. Schunck.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 21, May 26, 1879.

Refraction of Dark Heat.—P. DESAINS.—The author has never met with lenses so constructed as to collect to the same focus rays both from the beginning and the end of the dark spectrum. The study of the cold rays of the dark spectrum may lead to rules for constructing such lenses. If it is possible in the dark spectra to follow and recognise one and the same group of rays, then, in spite of the differences between the refractive and dispersive powers of the substances used, the absolute value of the refractions experienced by dark rays of one and the same wave-lengths in different diathermanous bodies may be determined.

Chemical Researches on the Formation of Coal.—E. FREMY.—The author holds that there are several kinds of isomeric cellulose, constituting the skeleton of plants. Coal is not an organised substance. The vegetal impressions presented by coal are produced as in shales or other mineral matters. The chief substances contained in the cells of plants under the double influence of heat and pressure produce bodies having a great analogy to coal. The pigments, the resins, and the fats of leaves if submitted to heat and pressure yield compounds which approximate to bitumens. The vegetable matter which gave rise to coal has undergone firstly the peaty fermentation, the coal being then formed by a secondary transformation.

Fluorescence of the Salts of the Earthy Metals.—J. L. SORET.—The author has already pointed out the beautiful violet fluorescence of solutions of cerium sulphate and chloride elicited only by the extreme ultra-violet rays of the induction-spark, the solar rays not being sufficiently refrangible for its production. He has since found that the solutions of many salts of the earthy metals possess analogous properties. He enumerates lanthanum chloride, didymium chloride and sulphate; terbium, yttrium, erbium, ytterbium chlorides; phosphoric chloride; thorium sulphate; zirconium sulphate and chloride; aluminium and glucinium chlorides.

Determination of the Lengths of Heat-waves.—M. Mouton.—Not susceptible of useful abstraction.

Diffusion of Lithia and its Presence in Sea-water.—E. Marchand.—The author shows that as early as 1850 he demonstrated the presence of lithia in sea-water, more than fifteen years before the spectroscopic researches of M. Bunsen, to whom M. Dieulafoy had in a recent paper ascribed the first discovery of this alkali in the sea.

Salts of Guanidin.—L. Joussetin.—The author merely describes methods of preparing certain of these salts.

Chemiker Zeitung.
No. 19, May 8, 1879.

Presence of Arsenic in Dark Water Colours.—Dr. H. Fleck.—Attention has been drawn to this subject by the sudden death of a mechanical draftsman. On a *post-mortem* examination the cause of death was first supposed to be an oxalate, and then a narcotic poison. Chemical investigation showed that the liver, kidneys, lungs, heart, and brain were impregnated with arsenic, though the oesophagus contained not a trace, and the stomach with its contents gave a barely perceptible arsenical mirror. The general circumstances of the case excluding the suspicions of suicide and of malicious poisoning, it was found that the deceased had been in the habit when drawing of drawing the pencil filled with colour between his lips in order to point it. The water-colours he had used were analysed, and whilst indian ink, gamboge, carmine, blue (2 which), red eosin ink, and neutral tint were found perfectly free from arsenic, a sample of sepia contained 3.08 per cent of arsenious acid, terra di Sienna 3.14, and a reddish brown colour, the name of which was indistinct, 3.15. Burnt Sienna, Vandyck brown, bistre, bladder green, brown ochre, indian red, umber, raw and burnt, were also found arseniferous. Most of these colours are essentially iron lakes. Hence it appears that the mere presence of ferric oxide, except in a hydrated state and accompanied by free magnesia in quantity sufficient to neutralise the acids of the stomach, does not act as an antidote to arsenious acid. This case seems likewise to prove that arsenic taken in minute doses can accumulate in the system until it can be readily recognised in all organs, and can exert a dangerous action. This result seems to prove that the impunity with which the peasants of Styria consume small doses of arsenic must depend upon circumstances not yet fully understood. (The exclusion of arsenic from such colours, in which it seems to play no essential part, should be insisted on by the authorities.)

A congress of German vine-growers will meet at Coblenz in September next, when the following questions will be discussed:—When and from what is sugar formed in grapes? Under what circumstances does sugar escape fermentation in wine? What recent results have been attained by the use of artificial manures in the cultivation of the vine?

Non-existence of Pentathionic Acid.—According to the *Bull. de l'Acad. Royale de Belgique*, Prof. Spring endeavours to prove that the supposed pentathionic acid is merely tetrathionic acid.

Ineradicable Ink.—The *Apotheker Zeitung* gives the following formula:—1.75 grms. aniline-black are ground up with 60 drops hydrochloric acid and 42 grms. alcohol, and the liquid is diluted with a hot solution of 2.5 grms. gum-arabic in 170 grms. water. If the aniline-black solution is diluted with a solution of 2.5 grms. shellac in 170 grms. spirit instead of gum-water, the result is an ink suitable for writing on wood, brass, or leather.

Preparation of Chemically Pure Tartaric Acid.—Ficius.—The decomposition of the calcium salt with sulphuric acid, or of the lead salt with sulphuretted hydrogen, does not yield a pure product. The author proposes to decompose the zinc salt with sulphuretted

hydrogen. Zinc tartrate is a scarcely soluble salt, easily obtained pure, even from very impure materials. The zinc sulphide obtained as by-product serves for the development of hydrogen sulphide, yielding a solution of zinc chloride, which is used for fresh precipitations of zinc tartrate.

Bulletin de la Société Chimique de Paris,
No. 8, April 20, 1879.

Preparation of Malonic Acid.—E. Grimaux and J. Tcherniak.—The authors first obtain cyanacetic acid and transform this into malonic acid by means of concentrated hydrochloric acid.

Formation of Aurin.—P. de Clermont and J. Frommel.—Phenol is converted into aurin by treatment with a mixture of carbonic oxide and oxygen gases under pressure at 250°. Neither carbonic oxide alone nor carbonic acid produces any effect.

Part Played by Auxiliary Acids in Etherification.—M. Berthelot.—Already noticed.

Certain Catalytic Phenomena due to Viscosity.—Antony Guyard.—The author considers that viscosity, like porosity, is capable of modifying chemical reactions, and possesses a true catalytic power. Glycerin, as a viscous substance, acts in two manners upon the reactions of metallic salts; it modifies the behaviour of these salts with known reagents because it is a non-volatile organic body, and especially because it is a viscous substance. Thus, if glycerin is mixed with a solution of chromic chloride, and ammoniac chloride and ammonia are added, the whole forms an emerald-green liquid, from which chromic oxide cannot be precipitated, just as if tartaric acid had been used instead of glycerin. Analogous reactions are manifested with some other metallic salts when mixed with glycerin and ammonia, or soda. When glycerin and the alkalis have no solvent power upon hydrated metallic oxides, precipitation takes place in the ordinary manner. If a large excess of glycerin is mixed with solutions of a double sulphate of potassa, of titanate acid, sulphate of alumina, ferric chloride, nitrate of lead, or stannous chloride and ammonia is added, all these oxides remain in solution, and are precipitated neither by boiling nor by the addition of water. But if hydrochloric acid is added enough to saturate at once the ammonia and the glycerin, and the acid is again supersaturated with ammonia, the oxides are precipitated as if the glycerin were not present. The addition of an alkaline chloride in sufficient excess to solutions of a metallic oxide in glycerin and caustic alkali determines its precipitation. The author has not succeeded in utilising these reactions in the quantitative separation of metallic oxides.

No. 9, May 5, 1879.

Compounds of Hydrogen Phosphide with Cuprous Chloride, and its Determination in Gaseous Mixtures.—J. Riban.—The reaction of these two bodies gives rise to several compounds, of which the chloride of cuproso-diphosphonium, $\text{Cu}_2\text{Cl}_2\text{PH}_3$, is well defined and crystallisable, though it soon undergoes decomposition on exposure to the air. Hydrogen phosphide when present in gaseous mixtures may be determined by agitation with a hydrochloric solution of cuprous chloride, in which it is rapidly absorbed.

Bromide of Tetraallylammonium and triallylamin.—H. Grosheintz.—The authors obtain this compound by passing a current of ammoniacal gas into an alcoholic solution of allyl-bromide. The mixture liberates heat spontaneously, and a crystalline mass is deposited. Triallylamin is prepared by distilling rapidly crude bromide of tetraallylammonium along with a large excess of recently-fused potassa.

New Organised Ferment of Urea.—P. Miquel.—The author has found in sewage an organism of the Bacilli

class, which, like *Micrococcus ureæ*, possesses the power of resolving urea into carbonate of ammonia, though quite distinct from the *Micrococcus* in its physical aspect. It appears, therefore, that two microscopical species possess approximately the same physiological function.

Precipitation of Lime by Alkaline Carbonates.—E. Drechsel.—The precipitation of calcic carbonate is complete in the cold in the course of fifteen minutes if aided by stirring. It may be effected indifferently either by sodic or ammoniac carbonates in the presence or absence of ammonia or sal-ammoniac. The calcic solution should be introduced into the alkaline carbonate, pouring in a small quantity of the former only at first and stirring for about five minutes before adding more. The solubility of calcic carbonate in alkaline liquids is very trifling.—*Journ. f. Prakt. Chemie.*

Gazzetta Chimica Italiana.
Anno ix., 1879. Fasc. iii.

Characteristic Reactions of Picrotoxin and of Certain of its Derivatives.—A. Ogliarolo.—If a small quantity of pure picrotoxin is dissolved in two drops of nitric acid at 14° and gently heated, there is obtained an amorphous residue of a reddish yellow colour. If two drops of potassa are added a fine bright red is obtained, which, on heating, passes to the colour of old blood. If 2 c.c. of a solution of picric acid at one-half per cent are mixed with potassa at 50 per cent in the cold nothing is observed but a yellow precipitate. On heating to a boil the precipitate dissolves, and the liquid is coloured orange. On cooling there are deposited small prisms of potassic picrate, and the liquid remains of a reddish yellow colour. If the experiment is repeated with the addition of picrotoxin, on boiling the colour of the solution becomes much deeper, and on cooling no crystalline deposit takes place, and the liquid remains intensely coloured. If a little picrotoxin is placed in a capsule and mixed with four or five drops of concentrated sulphuric acid, there appears a golden-yellow colouration which passes into a saffron-yellow; the picrotoxin dissolves, and on adding a little powder of potassic bichromate there is a violet-green colouration, and on dilution with water a clear solution of a yellowish green colour.

Distribution of Cerium, Lanthanum, and Didymium.—Prof. A. Cossa.—Already noticed.

Biedermann's Central-blatt.
Heft 4, April, 1879.

Chemico-agricultural Studies on a (Natural) Irrigation-meadow.—Prof. F. Ullik.—The manurial action of the water seems due to the dissolved substances, while the suspended matters are of little importance.

Ridge Cultivation.—Prof. E. Wollny.—The use of ridges or hillocks is advantageous only in retentive soils or in damp climates.

Nitrogenous Nutrition of Plants.—Prof. E. Heiden.—Cereals require an especial supply of nitrogen and repay it amply; the same may be said of potatoes. With leguminous plants this conclusion does not hold good. Rye and lupins cannot bear ammonia, at least in the form of sulphate, in the earlier stages of their growth.

Nitrogenous Constituents of Pasture-grass and Meadow-hay, and on their Digestion.—Dr. O. Kellner.—Of the total nitrogen present in grass nearly one-third is not in the condition of protein compounds, and can scarcely equal the carbohydrates in nutritive value. In hay the relative proportion of protein has greatly increased.

Can Rubidium Fulfil the Nutritive Function of Potassium in the Vegetable Cell?—Oskar Loew.—The author answers this question in the negative.

A New Oil Seed.—Dr. Eugen Wildt.—The seed in

question is that of *Lallemantia iberica*, cultivated in north-western Persia. It contains a smaller proportion of oil than do rape-seed, linseed, &c., whilst the residual press-cake is richer in nitrogenous matters.

Influence of Various Substances upon Crystallisable Sugar.—MM. H. Pellet and Durin.—A solution of sugar on standing and exposure to heat undergoes less change the stronger it is. Glucose converts cane-sugar into glucose, in proportion to the quantity of the former. In a solution saturated with cane-sugar this change does not take place. At certain temperatures mineral salts have a strong action upon cane-sugar, whilst the effect of organic salts is very small.

Treatment of Phosphatic Minerals.—T. Piltzer.—The phosphates are treated with sulphurous acid under pressure, which is produced by means of the carbonic acid liberated from the accompanying carbonates.

Absorption of Atmospheric Nitrogen by the Leguminosæ.—E. Gatellier has obtained good crops of lucern from an exhausted sandy soil where cereals proved a failure, and concludes that it has the power of fixing the nitrogen of the air. C. Marchand points out that lucern obtains the greater part of its nutriment from the subsoil, and that Gatellier's conclusion is therefore open to question.

Correspondenz-blatt des Vereines Analytischer Chemiker.
Vol. ii., No. 7, April 1, 1879.

Formula of Starch.—R. Sachsse.—Air-dried potato starch contains on an average 17.7 per cent of water. If this is supposed to be chemically combined, as may be inferred from the liberation of heat when it is taken up, starch would almost exactly represent the hydrate $C_{36}H_{62}O_{31} \cdot 12H_2O$.

Examination of Crude Pyrolignite of Lime.—A. Stromeyer.—5 grms. are distilled almost to dryness along with 50 c.c. of a solution of phosphoric acid, of sp. gr. 1.2, and the same volume of water. The operation is repeated with 50 c.c. of phosphoric acid, and again with the 50 c.c. of water. The acetic acid thus obtained is titrated with normal potassa.

Foreign Bitter Principle in Beer.—H. W. Langbeck.—The author prepared two samples of a fermented liquor from solution of glucose with small quantities of tartar, tartaric acid, kino, and a few drops of a mixture of formic and cœnanthic ether. Fermentation was set up by means of sound pressed yeast, and was maintained at a temperature of 18° to 20°. One sample, filtered through flannel after four days and allowed to stand for three weeks in a stoppered cask at 8°, yielded a pleasantly vinous liquid. The second sample, not filtered till after five days, tasted intensely bitter, and grew worse on standing. The newly-formed yeast, at first of a whitish yellow, had taken a brownish colour, died off, was precipitated by the more alcoholic character of the fluid, and formed with the alcohol in nascent state that substance which betrays itself by its bitterness in unhopped fermented liquors when the fermentation has been neglected. The compound in question is by no means innocuous. It was isolated by treating the liquor according to Dragendorff's methods I. and II. The author succeeded in obtaining it in a crystalline form and describes its reactions.

Determination of Solids in Milk.—H. Bering.—The author places about 0.09 grm. of calcined magnesia in a small platinum capsule, determines the tare, and introduces carefully, without touching the sides, an accurately weighed portion of the milk (from 1 to 2 grms.). On evaporation over a small open gas-flame, placed about 40 centimetres below the capsule, the sample is obtained perfectly dry in from two to three hours. That the dried mass is very feebly hygroscopic appears from the fact that 0.214 grm., after standing for twenty-four hours in a half wet room, had not gained 0.004.

Hannoversche Monatsschrift wider die Nahrungsfälscher.
Vol. ii., No. 2, February, 1879.

Sewerage, with Especial Reference to the Liernur System.—Dr. F. Fischer.—The author shows that in any town where the Liernur system was fully introduced, five-sixths of the urine would still find its way into the sewers. Even where cess-pools are lined with masonry, according to the observations made by Pettenkofer at Munich, nine-tenths of the excrementitious matters find their way into the subsoil. The cost of the Liernur system, including interest and sinking-fund for paying off initial outlay, is from 15s. to £1 yearly per head, and a system of sewers is still required.

Physical Properties of Fats.—The author does not consider that specific gravity affords a trustworthy means of deciding whether a sample of butter is genuine or sophisticated. Examination by polarised light supplies a much better characteristic. Mylius, who first called attention to this method, considers it of only limited applicability. The author is of a different opinion, as it is very rare for a butter to be sent for analysis after it has been melted. Pure butter, when examined with a magnifying power of 200 to 300 diameters, appears as a conglomerate of round and roundish drops of different sizes, interspersed here and there with characteristic salt-crystals. All melted fats after congealing take a crystalline structure. On examining a facitious butter we find not the above-mentioned globular drops, but more or less perfectly developed crystals, readily detected by the experienced eye, especially with an oblique illumination. All doubt is at once removed on examination by polarised light. The crystals come out very distinctly, and if the upper Nicol is slowly turned everything non-crystalline becomes gradually darker, whilst everything of a crystalline nature becomes lighter. The author finds, further, that different fats, like different minerals, produce characteristic differences in the polarisation colours. He announces the early publication of a series of plates showing the characteristic forms and colours of each fat, whether raw, melted, or crystallised from glycerin. Mutton tallow always gives a blue tone, and the contrasts when the Nicols are exactly crossed are sharper than in case of any other fat, except, perhaps, cacao-butter. The latter differs most characteristically from all other fats, and the play of colour from the deepest red to the brightest green does not admit of description. The fat of oxen displays merely green and white luminous effects. Small semilunar and vermicular bodies of a bright green appear by common light. Hog's lard displays many colours, especially red and blue, yellow, which is very conspicuous in cacao-butter, being wanting. These optical reactions are available for the detection of foreign fats fraudulently added to chocolate or cocoa.

MEETINGS FOR THE WEEK.

TUESDAY, 17th.—Zoological, 8.

WEDNESDAY, 18th.—Meteorological, 7.

THURSDAY, 19th.—Royal, 8.30.

— Philosophical Club, 6.30.

— Zoological, 8.

Chemical, 8. "On Gardenin," Dr. Stenhouse and Mr. Groves. "On the Action of Sulphuric Acid on the Hydrocarbons of the Formula $C_{10}H_{18}$," Drs. Armstrong and Tilden. "Researches on the Terpenes, Camphor, and Allied Compounds (Parts I. and II.)," Dr. Armstrong. "Contributions to the History of Starch and its Transformations," H. T. Brown. "On the Boiling-points of Certain Metals and Metallic Salts," Dr. Carnelly and W. C. Williams. "On the Determination of Nitric Acid by means of Indigo," R. Warington. "On Dry Copper-zinc Couples and Analogous Agents," Dr. Gladstone and Mr. Tribe. "Notes on the Purple of the Ancients," R. Schunck. Ballot.

ERRATUM.—P. 249, col. 2, line 6, for "invite" read "unite."

A German Chemist (Ph.D.), holding diplomas from University and Polytechnicum, who has already had experience in a manufactory, seeks a Situation in a Chemical Manufactory in England.—Address, K.D., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

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THE CHEMICAL NEWS.

Vol. XXXIX. No. 1021.

DETECTION OF FIRE-DAMP.

By Dr. R. ANGUS SMITH, F.R.S., &c.

THE detection of fire-damp, marsh-gas, or mine-gas, has occupied many minds since the time that Sir Humphry Davy began his enquiries and invented his celebrated precautions. I have often thought of two methods of testing for the gas, but only lately had them really tried. The first is by the use of the small instrument called, unfortunately, a compression syringe. It is commonly used for very rapidly compressing a small tubeful of air, so as to cause it to heat and ignite a piece of tinder. The tube used to be made of glass, but lately a method, inferior as I think, since the action becomes hidden, is employed, and the compression is made in a brass tube. A piston, which fits well, is put into the tube, and driven down rapidly by a blow with the palm of the hand on a smoothly rounded piece of wood at the top. As it was important to see the effect directly, a glass tube was necessary, and various substances were subjected to the heated air, so as to obtain a little experience as a beginning. There was no good tinder at hand, and cotton-wool in small quantities failed to ignite until it was well dried. Charcoal-dust ignited readily without special preparation. Minute quantities of gun-cotton at first did not take fire, and made the experimenters too bold, which ended in the shattering of the tube, which was 7 m.m. thick, and 6 m.m. internal diameter. When there was uncertainty the trial was made in the dark, and it was found that when a good lubricant was obtained there never was a failure; a spark was elicited readily. Coal-dust gave many sparks; ether fired readily, and a mixture of coal-gas and air exploded, bursting also more than one tube. It happened, however, that in trying the lubricant to see if the piston went down with ease, common air only being used, there was still a spark, and at last when the apparatus was in good order one was obtained on every occasion. It seemed clear that the olive oil used for lubrication was itself ignited. There were flashed alcohol, ether, turpentine, charcoal powder, coal powder, cotton-wool, olive oil, Young's lubricating oil.

The work, which had been at first difficult, had become at last too easy. The next step was to seek a non-combustible lubricant, and for a time a thick emulsion of oil and soft soap was used. This did not give any sparks alone, but the use of water is an objection. A strong glass tube, such as is sometimes sold for the compression, was shattered by one of the gas mixtures. For small explosions thinner tubes were used, $\frac{1}{4}$ m.m. in thickness of glass. Eventually this glass tube was put within a brass one 0.5 m.m. thick, in which was a window to allow the spark to be seen. Even this glass tube was shattered, giving way at the window, and bulging out the brass when 20 per cent of marsh-gas was added and exploded. I decided to have a much stronger tube, and that now made is unnecessarily thick. The glass is used only for about two inches at the bottom, and as the diameter must be at least as great as that of the brass it is necessary to put it in from below. To effect this a cap is made to screw upon the larger tube; this cap contains the glass tube, part of which forms the glass of the window. The screw is $\frac{1}{2}$ in. long, so as to hold well. The glass must not be closed below by the blowpipe, as a blow upon the conical base, or even a base of any shape, readily breaks it, and the piston may at times go quite to the bottom. The airtightness must therefore be produced by a good stuffing between the glass and the brass as well as

on the screw. The glass is covered with an india-rubber tube, and the two are put into the brass tube, previously having the bottom and sides well covered with glue, the best substance for this yet found. A mixture of resin and wax (and even caoutchouc if dry) was itself ignited by the improved mode to be mentioned.

This very strong apparatus has not yet been much tried, but before it was got a series of trials was made, by which it seemed certain that less than 5 per cent of marsh-gas could not be distinguished in air. In order to increase the delicacy of the process there was inserted a small amount of platinum-black, a very small amount is enough. The use of platinum I have had in view for several experiments in condensation of gases, and here it was found remarkably successful, bringing down the amount of marsh-gas to be detected to $2\frac{1}{2}$ per cent. On the other hand, it was necessary to use a lubricator without oil, so that the friction is considerably increased; at present we are using soap only.

The amount of gas detected is not minute, but it is a quantity which will not explode, or flash, or spark, unless under such exceptional circumstances as are here mentioned. It would be quite safe to make this explosion in any atmosphere, as the flash is confined to the tube, and a man could go in the dark and, as it were, feel the state of the air by making a spark as he went along. The apparatus needs nothing to be renewed except the air within. The piston must be taken out and care taken to bring in a fresh supply of air to be tried. The mode of doing this must have some attention. It must not be done by blowing as some might suppose, but it might be done by putting a fine tube into the cylinder and sucking out the air that had been compressed, and so letting fresh air in. It might also be done by having a stopcock below and letting the air in by that method, but the difficulty of keeping the joints tight is remarkable. The use of a loose plunger of wood to drive out the air is probably the easiest. The blow has a wonderful power. The cylinder and piston are only 8 m.m. in internal diameter, and the compression is more than thirty atmospheres, about 420 lbs. on the square inch.

Another method has gained my attention. Some time ago Mr. Ansell proposed to find the influx of fire-damp into the air of mines by using its diffusive power. By a well-known law of the diffusion of gases, the marsh-gas or fire-damp will pass into a porous vessel, if filled with common air; and if the covering is of caoutchouc the pressure will raise it. The rise of the cover may be used for making a communication with an electric bell, and this is the method Mr. Ansell used.*

The action is very rapid, but the raised caoutchouc soon falls. Besides this the air inside requires to be heavier than the outside gases which are expected to enter. It would be necessary to keep a supply of this air in order to make the apparatus self-acting, as it was intended. I did not obviate this difficulty, but designed simply the mode of filling the vessel with nitrogen, a gas nearly of the same specific gravity as air. When the experiment was to be made the vessel was filled with the nitrogen, and passed into the air to be tried. The pressure at once began within from the entrance of the lighter gas, and it raised a needle which was made to magnify the movement considerably.

This apparatus required, first, a solution of chloride of ammonium, next, chloride of lime. These mixed gave the nitrogen, which was made to pass into the vessel with the expandible cover. The pointers required some steady place on which to point, and the whole required a box. The plan was not found convenient, and it did not indicate under 5 per cent of marsh-gas as it was tried, but by varying the apparatus it could certainly be made to indicate much less. Such an apparatus may in some cases be found useful, and I give an account of it for that

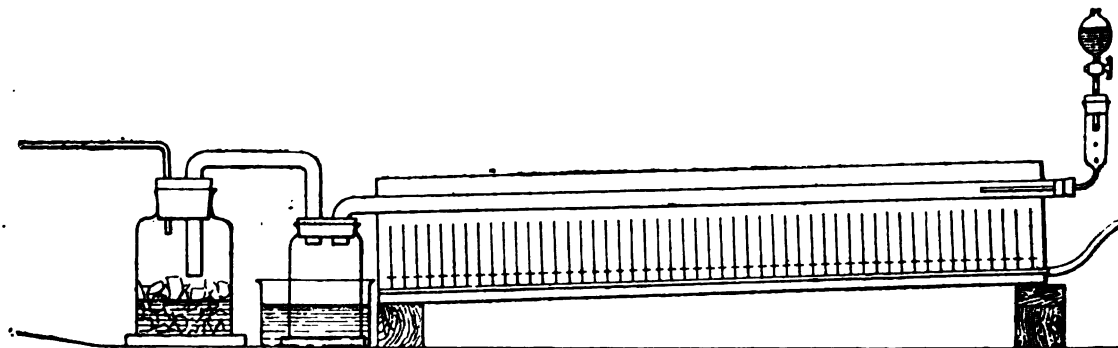
* A full description of the apparatus alluded to will be found in the CHEMICAL NEWS, vol. xii., p. 280, by Mr. George Frederick Ansell.

reason. The first-mentioned is certainly the handiest when people require to move about, and the experiment can be done in the dark; whereas this second plan needs a steady place and light. Still, it is sound in principle, and may be a useful supplement to Mr. Ansell's apparatus, which the late Lord Kinnaird, a lover of everything that promised good to the working classes, was very desirous of introducing as a common instrument in coal-mines.

READY METHOD FOR PREPARING DIPHENYL.

By WATSON SMITH, F.C.S., F.I.C.

By the adoption of the following method I have found that a rich yield may be obtained of the above hydrocarbon, and the experiment is easily carried out, proceeds rapidly, and without any danger. 62.4 grms. of benzene are mixed in a flask with 52 grms. of tin tetrachloride, or at least this proportion is observed, and the mixture is poured little by little into a dropping-tube, which allows it to fall drop by drop into a tube of Bohemian glass, itself lying in a combustion-furnace, and maintained at a bright red heat. The combustion-tube should be bent at the outlet, almost at right angles, so as to pass through a cork fitting into a wide-necked bottle. A tube passes from this bottle to another large bottle containing some water and also a quantity of blotting- or filter-paper. This arrangement serves to partially condense the hydrochloric acid evolved,



but another tube passes from this bottle into a draught-place, or through a window, to lead off the uncondensed gas. There is no need to pass the evolved gases through a Liebig's condenser, for if a bright red heat be maintained not more than one drop of undecomposed benzene would be condensed, the above weights being taken. The diphenyl is found mixed with stannous chloride in the receiver in solid masses. Separation is easily effected by warm concentrated hydrochloric acid, which dissolves the stannous chloride, and leaves behind diphenyl. The latter is purified by distillation alone, and finally with steam.

In a paper read before the Chemical Society, and published in the *Journal* (July, 1876, and November, 1877), I recommended distilling the contents of two flasks into the red-hot tube, one containing the tin tetrachloride, the other the benzene, and so that the vapours of the benzene passed through the flask containing the tin tetrachloride heated up to boiling. I now find that the method above recommended is much safer and easier of execution.

Widmannstätten's Figures on Artificial Iron.—J. Lawrence Smith.—These figures, developed by the action of an acid upon a polished surface, have been considered characteristic of meteoric irons, some of which, however, do not show these figures. The author has succeeded in producing them upon silicide of iron. *Comptes Rendus*.

ON NITRIC NITROGEN IN GUANO.*

By ROBERT R. TATLOCK, F.R.S.E., F.C.S.

PART I.

FROM the results of some experiments which I made many years ago, I was led to the conclusion that a large proportion—in some cases nearly the whole—of the nitrogen existing as nitrates in guano appears, and is estimated as ammonia, in the ordinary process of combustion with soda-lime. As there are no means known, so far as I am aware, even up to the present time, of preventing the conversion, in presence of organic matter, of a very indefinite but always large proportion of the nitric nitrogen into ammonia, it follows that all determinations of the latter made in guano are inaccurate unless they are based upon the complete decomposition of the nitrates with production of ammonia from the whole of their nitrogen; and I am not aware that up till now this has been accomplished, or even the necessity for it admitted. The amount of error will depend chiefly on the ratio of the nitrates to the organic matter present, but will also be much influenced by the nature of the latter as well as by the completeness or otherwise of its admixture with the subject of the analysis. As it is not uncommon for natural guanos to contain from 1 to 2 per cent of nitrates, the error due to partial or imperfect decomposition of the latter is considerable, but in the case of guanos or artificial manures which have been purposely mixed with a considerable proportion of a nitrate, such as nitrate of

soda, it is much augmented, and if, say 10 per cent of the last named substance has been employed, the error may and probably will be as great as 1 per cent of ammonia—to the low side if it be assumed that the whole of the nitric nitrogen is obtained as ammonia, and to the high side if it be assumed that none of is obtained, in that form—in which case it is of course determined by a separate process and added to the result obtained by the ordinary combustion with soda-lime. I was surprised to learn, recently, that the latter was the practice of chemists who had had a very large and varied experience in agricultural analyses, and the effect of it is that purchasers of guanos and manures containing nitrates either naturally, or artificially introduced, on the basis of their analyses, must pay for a large proportion, and in many instances practically the whole of the nitric nitrogen twice over. It is possible that this practice, erroneous as it is, has arisen from the circumstance that when nitrates are heated to redness with soda-lime, no ammonia is obtained; at any rate I have ascertained this by direct experiment. The result is very different, however, when organic matter is present, ammonia being always produced, varying in amount according to the nature and proportion of the organic substance and other circumstances. Being desirous of obtaining some reliable method by which the whole of the nitrates could be converted into ammonia during the

* A Paper read before the Newcastle-upon-Tyne Chemical Society March 27, 1879.

Organic Substance used.	Quantity of Organic Substance used. Grammes.	Quantity of KNO ₃ used. Grammes.	Ratio of Organic Substance to KNO ₃ .	Weight of (NH ₄) ₂ Pt. Cl ₂ obtained.	Percentage of Nitric Nitrogen obtained.
Starch	0.6	—	—	0.010	—
Do.	0.6	0.05	12 to 1	0.054	39.88
Do.	0.6	0.2	3—1	0.210	45.30
Do.	2.0	0.1	20—1	0.112	50.74
Camphor	0.6	—	—	—	—
Do.	0.6	0.2	3—1	0.121	26.38
Wood charcoal .. .	0.3	0.2	1½—1	0.051	11.56
Albumen (white of egg) ..	0.6	—	—	1.276	—
Do.	0.6	0.1	6—1	1.386	49.94
Sugar (candy) .. .	2.0	—	—	—	—
Do.	0.6	0.1	6—1	0.140	63.35
Do.	0.6	0.1	6—1	0.150	67.92
Do.	1.0	0.1	10—1	0.178	80.63
Do.	1.5	0.1	15—1	0.191	86.53
Do.	2.0	0.1	20—1	0.190	86.07
Do.	2.0	0.1	20—1	0.150	86.07
Do.	3.0	0.1	30—1	0.215	97.40
Sugar	0.8	—	—	—	—
Camphor	0.2	0.1	10—1	0.137	61.98

combustion, and as it seemed hopeless to expect to find any means of preventing the production of ammonia, to a large extent, from them, in presence of organic matters such as are invariably present in guano, I undertook the following series of experiments with the object of determining the effect of organic matters of different kinds and in different proportions in the production of ammonia from alkaline nitrate, in the ordinary combustion process by soda-lime.

In the trials made with starch the necessary correction was made for the organic nitrogen naturally present (which was found to be 0.10 per cent) before the results were recorded in the table.

The combustions were made in the ordinary way, 40 grammes of soda-lime being employed in each case, the ammonia collected in dilute hydrochloric acid contained in nitrogen bulbs, as usual, and weighed as chloro-platinate of ammonium. The organic matter was intimately ground with the nitrates; this mixture was then first ground with a portion of the soda-lime, and the result well mixed with the remainder. (See above.)

It will be seen from this table that the following conclusions are deducible from these experiments:—

1. That using 3 of organic matter (starch) to 1 of nitrate, 45.30 per cent of the nitric nitrogen can be obtained.
2. That in no case was the whole of the nitric nitrogen converted into ammonia, the greatest proportion being 97.40 per cent.
3. That the results are somewhat variable even with the same proportion of the ingredients, something always depending on the completeness with which the mixture is made.

It will readily be conceded that in guanos where the proportion of organic matter to nitrates is very high, as is the case in ammoniacal guanos, it is quite erroneous to determine nitric nitrogen by a separate process and add it to the nitrogen obtained by combustion with soda-lime, it being already included therein. It follows from this consideration and from these experiments that it is best to conduct the combustion in such a way as to convert as much as possible of the nitric nitrogen into ammonia, and that if it is determined separately in order to distinguish it from that existing in other forms, it should not in these circumstances be added to the nitrogen, in which it is already included.

The following detailed analysis which I have made of a sample of "Huanillos" guano will serve to show, by observing the ratio of the organic matter to the nitrate, that at least one-half, probably much more, of the nitrogen contained in the nitrate will be estimated in the process of combustion:—

	Per Cent.
SOLUBLE IN WATER	100.00
Phosphate of ammonia .. .	14.29
Phos. mag. and ammonia .. .	1.30
Chloride of ammonium .. .	2.38
Sulphate of potash .. .	6.42
Sulphate of soda .. .	4.81
Nitrate of soda .. .	1.55
Chloride of sodium .. .	5.74
INSOLUBLE IN WATER.	
Phosphate of lime (tribasic) .. .	14.91
Carbonate of lime .. .	3.77
Carbonate of magnesia .. .	2.64
Sand .. .	5.08
Organic matter .. .	21.34
Water .. .	15.77
Soluble phosphoric acid .. .	7.11
Equal to tribasic phosphate of lime .. .	15.33
Insoluble phosphoric acid .. .	6.83
Equal to tribasic of phosphate lime .. .	14.91
Total phosphoric acid .. .	13.94
Equal to tribasic phosphate of lime .. .	30.44
Nitrogen in ammoniacal salts .. .	5.83
Equal to ammonia .. .	7.08
Nitrogen in organic matter (potential) .. .	1.43
Equal to ammonia .. .	1.73
Nitric nitrogen (in nitrate) .. .	0.25
Equal to ammonia .. .	0.31
Total nitrogen .. .	7.51
Equal to ammonia .. .	9.12

With regard to the best methods for the determination of the nitric nitrogen in guanos there is still a great difference of opinion. Methods based upon the liberation of nitric acid, and the determination, directly, of the oxidising power of the liberated acid, are erroneous and misleading, from the fact of the well-known action of nitric acid, especially when largely mixed with strong sulphuric acid upon soluble organic matters such as are always present in ammoniacal guanos. The indigo method of Boussingault is an example of these, although it is recommended by experts in agricultural analyses; and that of Harcourt, by distilling with zinc, iron, and caustic alkali, although a minutely accurate one for estimating nitrogen, when properly applied, as I can testify from a long experience of it, is not applicable to guano on account of the soluble nitrogenous matters always present, and which even permanganate of potassium only partially destroys. Yet this method is recommended by skilled analysts, with the sole precaution of removing the ammoniacal salts before applying it,

although the inevitable result of its use is that the organic nitrogen is included with the nitric nitrogen. The consequence of employing this method in conjunction with the practice of adding the nitric nitrogen to that obtained in the combustion process is that part of the nitrogen is stated, and consequently paid for three times, the organic nitrogen being inadvertently determined with the nitric nitrogen, and both added to the nitrogen obtained by combustion, which already contains both.

Pelouze's method, based upon the oxidation of ferrous salts by the liberated nitric acid, in an atmosphere of carbonic acid, I have found to give good results in some cases; but they are no doubt too low where much soluble organic matter is present, even if the latter be removed as far as possible by permanganate of potassium.

Crum's process (*Proceedings of the Philosophical Society of Glasgow*, 1848, p. 162), besides being a really scientific one, and capable of great accuracy, is free from the objection of giving too low results in presence of soluble organic matter; for although the latter is acted upon by the nitric acid, it is a matter of indifference what reduces the latter if only nitric oxide be produced, by the measurement of which the nitric nitrogen is estimated. It seems necessary, however, to prove, where organic matter is present, that the evolved gas consists entirely of nitric oxide, as a little nitrogen sometimes appears, probably as a result of the action of the strong sulphuric acid on nitrogenous organic matter. This can easily be done, as Crum describes, by introducing into the nitrometer some warm solution of iron sulphate, observing the amount of gas absorbed and regarding this only as nitric oxide.

I am at present endeavouring to obtain the perfect resolution of the nitric nitrogen into ammonia, and am also conducting some experiments with the view of finding the method best adapted for estimating nitric nitrogen in guanos and manures.

ON THE DIFFERENT ANILINE-BLACKS.*

By JUSTUS WOLFF.

IN the first instance there are two different series of aniline-blacks, viz., those which are produced in or on the fibre, and those which are manufactured first and then applied to the fibre by the usual dyeing process.

The first-named series were invented by the late Mr. John Lightfoot, of Accrington, in the year 1866, and are extremely well adapted for printing black on vegetable tissues, being in some cases the best black-print out.

Many various fruitless and expensive efforts have been made to dye fibres and fabrics by the Lightfoot process and its improvements, but have failed, although the shades of the black produced by some of the methods in these series are extremely beautiful.

The greatest difficulty in dyeing according to Lightfoot's process is to get the shades evenly distributed over the fibres and fabrics, and it has not yet been overcome satisfactorily, although many experiments on a very large scale have been made for that purpose in Bradford and Manchester dye-works with cotton goods.

The difficulties in dyeing worsted and silk tissues by these and kindred processes in even shades are still greater, and that is the reason why this beautiful and fast dye is not produced on yarns and fabrics on a large scale except by printing on cotton, in which application it far surpasses all other black print by beauty of shade.

The methods of dyeing yarns and woven fabrics according to the Lightfoot process consist principally in soaking them thoroughly and evenly in a strong solution of aniline hydrochloride, with or without free aniline, and potassium chlorate, with or without the addition of other—especially metallic—compounds, afterwards exposing the goods

in a moderately warm place to the action of air for such a length of time until all is of a dark green colour, and subsequently passing them through a warm bath of soda, which develops the black in a short time. Instead of the soda-bath, a bath with small quantities of chrome and hydrochloric acid develops a much deeper and finer black, which does not turn green.

Of course there are nearly as many modifications of this method of dyeing as there are dyers, colourists, and chemists, who have been trying it, and have modified the original process with a view to improve upon it, many of them succeeding to a certain extent, but none completely.

The introduction of vanadium in these processes (which is not new, as the late Mr. John Lightfoot mentioned it in his Patent specifications) was certainly a great improvement, but even with its aid it was impossible to overcome the difficulties in dyeing mentioned above. Even supposing these difficulties had been overcome, the process of dyeing Lightfoot-black would be a very tedious one on account of the length of time needed for the development of the green colouration and the great number of baths the goods would have to pass.

The Lightfoot-blacks can be divided into those which turn green by exposure to air, and those which remain black under the same circumstances. The first are common and the second the oxidised Lightfoot-black. The shades of these series of blacks run from blue, dark bluish-grey, blue-black, grey, and black to brown-grey brown-black, and even to brown and black-brown.

The first link of these series is the blue which the Dr. Crace-Calvert invented and obtained by the action of aniline hydrochloride with a smaller quantity of potassium chlorate than for black, and some ferrous sulphate in order to moderate the oxidation. Next to this blue ranges Lightfoot's blue-black, and then follow the other shades in the succession given above.

These blacks, in common with the other aniline-blacks, are mixtures of at least two shades, viz., of a very deep blue and deep brown, the latter varying from reddish brown to yellowish brown, which quality enables it to produce many different shades of black in combination with the deep blue.

The purer the aniline—that is, the less toluydin it contains—the bluer is the black produced by that process, thus showing that the brown colouring matter accompanying the blue in the Lightfoot-blacks owes its origin to toluydin contained in the commercial anilines, even if only in small quantities.

Some metallic compounds—as, for example, of copper, cerium, vanadium, and certain others—have the property to deepen this dark blue to a very fine blue-black, even if employed only in small quantities, by their ability to increase the strength or power of oxidation.

The stronger the action of oxidation, up to a certain degree, on the aniline or the aniline salt the darker becomes the product of that oxidation, whilst the weaker the action of oxidation the bluer the shade obtained. On this principle a blue or black can be produced from the same quality of aniline.

As to the chemical constitution of the Lightfoot-blacks not much is known, and the investigations made to that purpose have not yet thrown much light on that dark object.

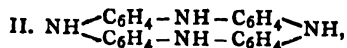
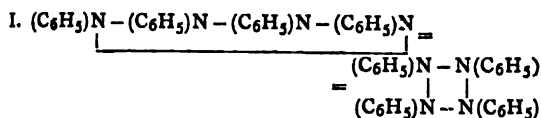
According to Reinbeck's results of research, the Lightfoot-black is a powerful base of violet-black colour, forming with acids green coloured compounds.

A. Müller's formula of Lightfoot-black, as the result of direct elementary analysis, $C_{12}H_{14}N_2O_{11}$, is an improbable one on account of the large proportions of hydrogen and oxygen. Without regarding this last-named defect, the proportion of carbon and nitrogen would allow us to consider it as a derivative from diphenylen-diamin by powerful oxidation.

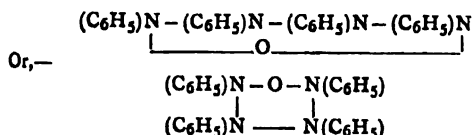
A more reliable elementary analysis published by Goppelsröder (*Dingler's Polytechnisches Journal*, ccxv., 439 to 448) leads to the empyric formula $C_{24}H_{26}N_4$ for the

* All temperatures are given in Centigrades.

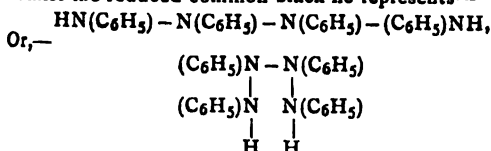
common Lightfoot-black; he interprets in it the following ways.—



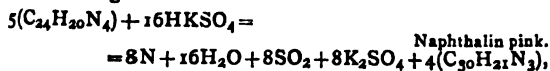
of which he considers the first as the most probable one. The chemical constitution of the oxidised black is, then,—



Whilst the reduced common black he represents—



With potassium bisulphate he produces naphthalin-pink from the Lightfoot-black:—



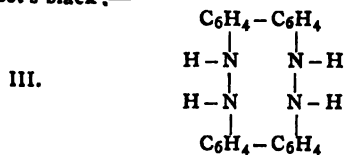
wherein the following decomposition of potassium bisulphate takes place:—



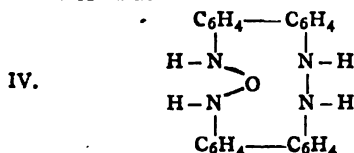
Another author produces by treatment of Lightfoot's black with aniline a fine aniline-pink of the formula—
 $\text{C}_{36}\text{H}_{33}\text{N}_5$.

These investigations have been made on the aniline-black produced by electrolysis, which is identical with the Lightfoot-black. All these formulæ of aniline-blacks show that they are the products of a powerful oxidation simultaneously with a considerable condensation.

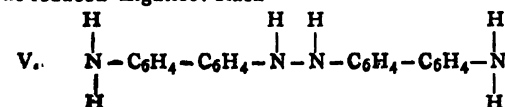
It is very probable that in Lightfoot-black a connection of nuclei has taken place (an interpretation supported by the production of naphthalin-pink with bisulphate of potassium, as stated by Goppelsröder, by the property to form products of substitution with aniline,—aniline-pink—and also by some analogies coming forth in the course of this paper), which leads to the following formulæ for Lightfoot's black:—



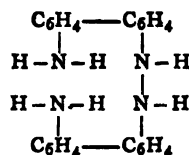
The oxidised black—



The reduced Lightfoot-black—



Or—



Turning now to the other aniline-blacks, viz., those which are manufactured from aniline first, and then when ready are applied to the fibres by the usual dyeing process, we meet with two series, which we will keep distinct by the names used for them in commerce, viz., indulin and nigrosin.

The name nigrosin I gave to that series of black which I invented in the year 1862, in the laboratory of Messrs. Appold Frères, in Sulzbach, near Saarbrücken. But since then this name has been usurped by some firms for colouring matters belonging to the indulin series in order to sell an inferior article at the price of the real nigrosin.

The first link of the indulin series I found out in 1865 by treating the bases of magenta-refuse with aniline and acetic acid, a process very similar to that of the production of aniline-blue from magenta. The spirit-soluble indulin produced by that process was converted with oil of vitriol into the water-soluble indulin, fraudulently called by some firms nigrosin.

The indulins are manufactured in several ways; first from magenta refuse:—

The refuse is treated with boiling water containing hydrochloric acid in order to extract the salts of mauvanilin, rosanilin, and chrysianilin completely, and to leave the violanilin salt undissolved, which is decomposed with caustic soda, and by this treatment the violanilin base is obtained in a free state. To convert this violanilin (or also mixtures of bases containing it in the largest proportion), a mixture of say, 10 parts dry violanilin, 6 parts commercial acetic acid (of the equivalent 120), and 20 parts aniline for blue, is heated to between 140° to 160° as long as ammonia is developed, and till the mass dissolves in alcohol (acidulated with acetic acid) in the desired shade. Arrived at that state, the excess of aniline is liberated by addition of caustic soda sufficient to neutralise the 6 parts of acetic acid (which are in the mixture), and driven off by a current of steam. The indulin base thus obtained is separated from the solution containing acetate of soda, washed, powdered, and dried, and may be used in that state directly for conversion into the water-soluble state; but it may also be purified by treating it several times with boiling acidulated water in order to obtain the salts of the indulin bases, which also have to be dried before conversion. This conversion is effected in the following manner:—

Into three or four parts of oil of vitriol of 66° B., or, if needed, of fuming oil of vitriol (Nordhausen vitriol oil), or mixtures of both, heated up to about 100°, one part of dry indulin base or its salts is given in slowly whilst the liquid is well agitated. When all is in, the heat is raised to about 120° to 140°, and kept there for such a length of time (about five hours) till a sample drawn and completely washed with water (in order to remove all free sulphuric acid), and treated with ammonia at about 60° or 70°, dissolves quickly and completely. This point reached, the solution of the colour in oil of vitriol is, whilst yet hot, poured into about five times its weight of clear cold water, which is well agitated, then settled for several hours, the water drawn off, and the sediment repeatedly washed in this way with fresh quantities of water till it produces no perceivable sour taste. Then it is filtered and boiled with caustic soda solution, just sufficient to dissolve it and to form a neutral salt, and then dried down at a temperature not higher than 70°. Sometimes in place of soda ammonia is used. It forms then the water-soluble indulin.

Another way to produce indulin is by heating a mixture of 10 parts aniline (possibly pure preferred) with 20 parts of syrupy arsenic acid (containing 70 per cent of dry

acid) to 185° to 190° for such a length of time, till it forms after cooling a dull yellowish, bronze-coloured, brittle substance.

At that state most of the aniline is converted into violanilin, which is obtained pure by neutralisation of the arsenious and arsenic acid in the melt with caustic soda, driving off the un-converted aniline by a current of steam, separation of the melted violanilin base from the water (formed by condensation of steam), powdering, and drying it. This base is converted with aniline and acetic acid into the spirit-soluble indulin or its base (as described above), which is made soluble with oil of vitriol, similarly as the indulin base from magenta refuse.

Violanilin is also produced by the action of arsenic acid on aniline hydrochloride, or other suitable salts of pure aniline; further, by the action of pure nitro-benzene on pure aniline alone, or mixed with aniline salt, and in general by the action of suitable oxidising or dehydrogenating substances on pure aniline or suitable aniline salts at a temperature of 185° to 190°. Also, for example, by a current of chlorine gas into aniline at 180° to 190°, and then, on the same principle, with a mixture of nitrate of aniline with aniline salt, or by a mixture of nitrate of soda with aniline salt in excess, and so on.

Of course in all these processes some other substances besides violanilin are coming forth, but the latter will be the principal one if the operation is conducted very carefully, and the temperature of forming violanilin does not rise higher than 190°. Above that temperature other decompositions take place, and by-products will be formed to a great extent—as, for example, triphenylen-diamin-blue, &c.—but which to take into consideration would complicate this paper to an enormous extent. Therefore I will only treat on the principal substances formed in the largest proportions by these processes, under the supposition that they are carefully conducted.

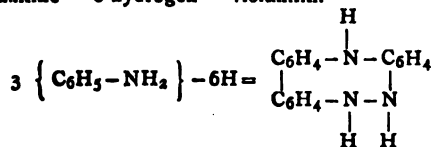
As an explanation of the chemical reactions going on in the processes above described I give the following:—

Violanilin is the product of elimination of hydrogen from pure aniline, and simultaneous condensation.

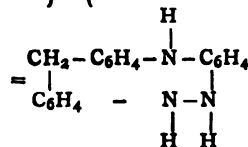
Mauvanilin is the product of elimination of hydrogen and simultaneous condensation of a mixture of two molecules of aniline and one molecule of toluidin.

Both these reactions are represented in the following:—

3 aniline - 6 hydrogen = violanilin.

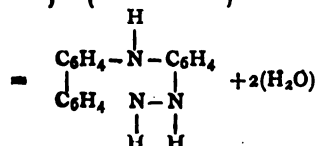
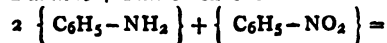


2 aniline + 1 toluidin - 6 hydrogen = mauvanilin.

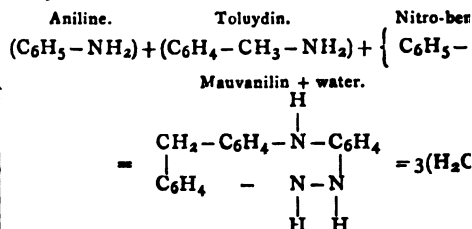


The reaction of nitro-benzene on aniline, or aniline hydrochloride, or mixtures of both, is as represented by the following formulæ:—

2 aniline + 1 nitro-benzene = 1 violanilin + 2 water.



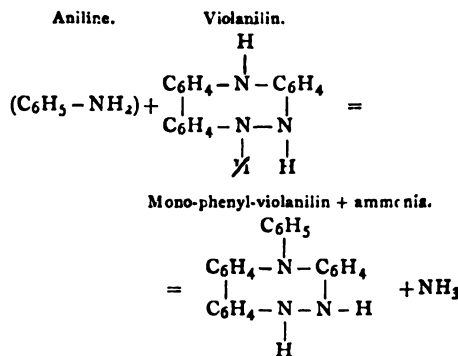
And that of nitro-benzene on a mixture of aniline and toluidin—



If there is no hydrochloride of aniline, or of that of aniline and toluidin, and no free aniline or toluidin to be acted upon, then the process must be watched carefully, and by all means be kept below 190°, else a quantity of by-products is formed, much easier than the same process without hydrochloric acid. A sudden reaction (especially on a large scale) takes place, producing a rise of temperature by itself at once, and forming a large quantity of by-products. Therefore, if the temperature should rise above 190° the mixture should be cooled down by the addition of water whilst agitated, and by these means the temperature should be kept within the limits given above.

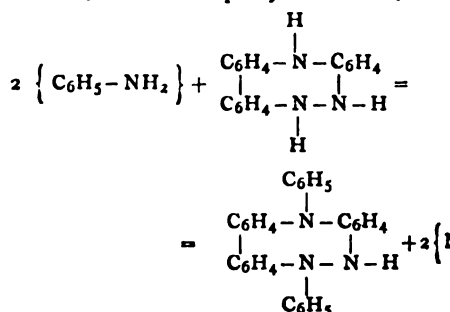
The conversion of violanilin with or without mauvanilin into spirit-soluble indulin is caused by the substitution of one, two, or three hydrogen atoms in the appendants by phenyl.

If acetate of violanilin with or without mauvanilin is heated with aniline for blue to 140° to 160°, then indulin is formed, and the above-named substitution takes place, producing in the first instance the mono-phenyl-violanilin.



In the second stage of the process diphenyl-violanilin is formed—

2 aniline + violanilin = diphenyl-violanilin + 2 ammonia.



If that process is carried on so far till the formation of ammonia ceases, then a complete substitution of the hydrogens in the appendants has taken place, and triphenyl-violanilin has been formed.

acid) to 185° to 190° for such a length of time, till it forms after cooling a dull yellowish, bronze-coloured, brittle substance.

At that state most of the aniline is converted into violanilin, which is obtained pure by neutralisation of the arsenious and arsenic acid in the melt with caustic soda, driving off the un-converted aniline by a current of steam, separation of the melted violanilin base from the water (formed by condensation of steam), powdering, and drying it. This base is converted with aniline and acetic acid into the spirit-soluble indulin or its base (as described above), which is made soluble with oil of vitriol, similarly as the indulin base from magenta refuse.

Violanilin is also produced by the action of arsenic acid on aniline hydrochloride, or other suitable salts of pure aniline; further, by the action of pure nitro-benzene on pure aniline alone, or mixed with aniline salt, and in general by the action of suitable oxidising or dehydrogenating substances on pure aniline or suitable aniline salts at a temperature of 185° to 190°. Also, for example, by a current of chlorine gas into aniline at 180° to 190°, and then, on the same principle, with a mixture of nitrate of aniline with aniline salt, or by a mixture of nitrate of soda with aniline salt in excess, and so on.

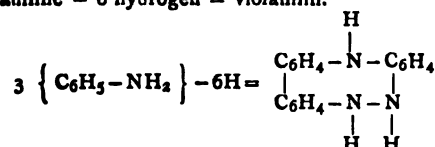
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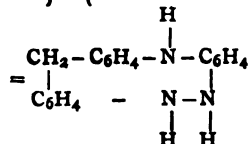
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Both these reactions are represented in the following:—
3 aniline - 6 hydrogen = violanilin.

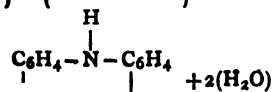
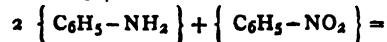


2 aniline + 1 toluidin - 6 hydrogen = mauvanilin.

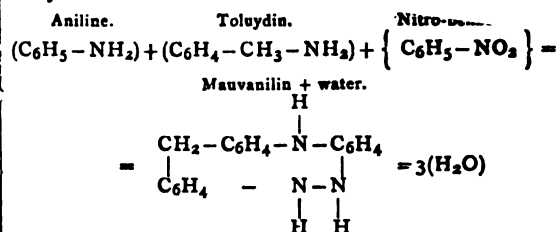


The reaction of nitro-benzene on aniline, or aniline hydrochloride, or mixtures of both, is as represented by the following formulæ:—

2 aniline + 1 nitro-benzene = 1 violanilin + 2 water.



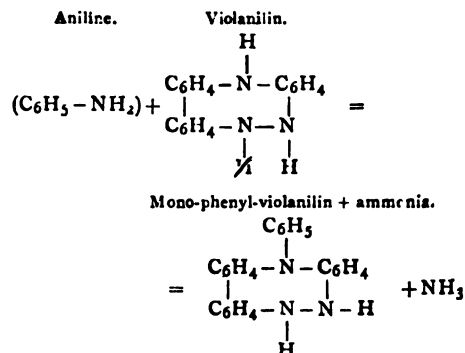
And that ...
toluidin—



If there is no hydrochloride of aniline, or of the mixture of aniline and toluidin, and no free aniline or toluidin to be acted upon, then the process must be watched very carefully, and by all means be kept below 190°, else a large quantity of by-products is formed, much easier than in the same process without hydrochloric acid. Even a sudden reaction (especially on a large scale) may take place, producing a rise of temperature by itself up to 140° at once, and forming a large quantity of several by-products. Therefore, if the temperature should rise above 190° the mixture should be cooled down by a thin stream of water whilst agitated, and by these means the temperature should be kept within the limits given above.

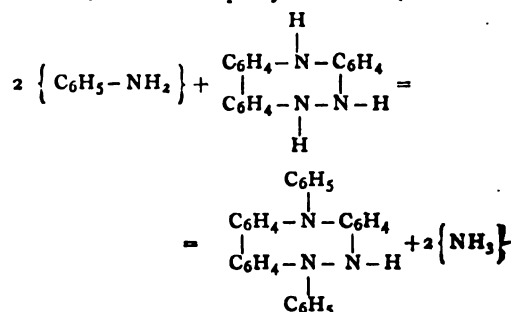
The conversion of violanilin with or without mauvanilin into spirit-soluble indulin is caused by the substitution of one, two, or three hydrogen atoms in the violanilin appendant by phenyl.

If acetate of violanilin with or without mauvanilin heated with aniline for blue to 140° to 160°, then ammonia is formed, and the above-named substitution takes place, producing in the first instance the mono-phenyl-violanilin



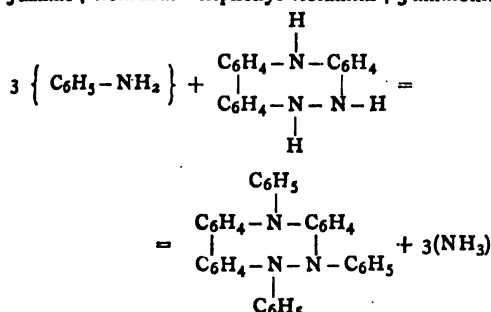
In the second stage of the process diphenyl-violanilin is formed—

2 aniline + violanilin = diphenyl-violanilin + 2 ammonia



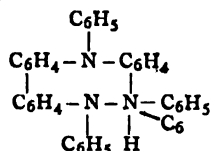
If that process is carried on so far till the formation of ammonia ceases, then a complete substitution of hydrogens in the appendants has taken place, and

3 aniline + violanilin = triphenyl-violanilin + 3 ammonia.



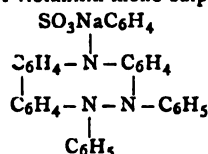
In the most spirit-soluble indulins the triphenyl-violanilin is predominant in quantity; in many of them the mono- and diphenyl-violanilin and mauvanilins accompany it.

Thus, indulin will be principally triphenyl-violanilin hydrochloride :—

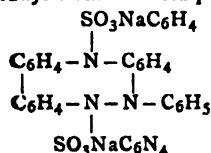


By the treatment of these bases and their salts with oil of vitriol, as described above, the four conjugated acids and their salts may be produced.

Sodium triphenyl-violanilin-mono-sulphonate—



Sodium triphenyl-violanilin-disulphonate—



And also the tri- and tetra-sulphonates. And, of course, besides these the corresponding sulphonates of the accompanying compounds are formed simultaneously. The mono-sulphonic acids are insoluble in water, but their alkali salts are easily soluble. The disulphonic acids are only sparingly, but their alkali salts are easily, soluble.

The tri- and tetra-sulphonic acids, and their salts of the phenylated viol- and mauvanilins, are soluble in water, the last the most, and these form the principal constituents of the commercial water-soluble indulins, sometimes containing some of the nigrosin sulphonic acids or their salts.

The properties of the spirit- and water-soluble indulins are very nearly like those of the aniline-blues, and their constitutional formulæ are also much alike.

The spirit-soluble indulin dyes wool, silk, cotton, and other fibres grey in various shades, whilst it is difficult to dye black shades with. It is dissolved in acidulated alcohol (or methylated spirit) by boiling. The filtered solution is added to the acidulated cold bath, in which the goods to be dyed are immersed, and which is heated up slowly to the boiling-point, and kept there till the desired shade is reached.

The spirit-soluble indulin dissolves at about 115° in 2 to 3 parts of its weight of glycerin containing 5 per cen.

hydrochloric acid. After it is dissolved water is added slowly under agitation to five times the weight of glycerin, by which means a solution is obtained which can be used for dyeing. But all solutions of spirit-soluble indulin have the disadvantage that the indulin separates quickly in the bath, rendering the dyed shades uneven.

The water-soluble indulins are dissolved in warm water, and the solution given in the luke-warm acidulated bath where the goods had been immersed before, and then the bath is heated nearly up to the boiling-point, and kept there till the desired shades are produced. The water-soluble indulins dye good shades of light and dark greys, even approaching black, which shade to reach fully offers some difficulties, so that they are scarcely used for black-dyeing on a large scale; besides, the black, when reached, is of a dull and inferior shade, and only slightly resists light, air, and soap.

(To be continued.)

ELECTRIC LIGHTING.

THE Report of the Select Committee of the House of Commons on Electric Lighting was issued June 18th. It is as follows :—

The general nature of the electric light has been well explained in the evidence of Prof. Tyndall, Sir William Thomson, Dr. Siemens, Dr. Hopkinson, and others. It is an evolution of scientific discovery which has been in active progress during the whole of this century. Essentially the electric light is produced by the transformation of energy either through chemical or mechanical means. The energy may be derived from a natural force, as, for instance, a waterfall, or through combustion of a material in the cells of a voltaic battery, or of fuel in a furnace. The energy being converted into an electric current, may be used to manifest electric light by passing between carbon points, or by rendering incandescent solid bodies, such as iridium. A remarkable feature of the electric light is, that it produces a transformation of energy in a singularly complete manner. Thus the energy of one-horse power may be converted into gaslight, and yields a luminosity equal to 12-candle power. But the same amount of energy transformed into electric light produces 1600-candle power. It is not therefore surprising that while many practical witnesses see serious difficulties in the speedy adaptation of the electric light to useful purposes of illumination, the scientific witnesses see in this economy of force the means of great industrial development, and believe that in the future it is destined to take a leading part in public and private illumination. There is one point on which all witnesses concurred, that its use would produce little of that vitiated air which is largely formed by the products of combustion of ordinary illuminants.

Scientific witnesses also considered that in the future the electric current might be extensively used to transmit power as well as light to considerable distances, so that the power applied to mechanical purposes during the day might be made available for light during the night. The Committee only mention these opinions as showing the importance of allowing full development to a practical application of electricity, which is believed by competent witnesses to have future important bearings on industry.

So far as the practical application of the electric light has already gone, there seems to be no reason to doubt that it has established itself for lighthouse illumination, and is fitted to illumine large symmetrical places, such as squares, public halls, railway stations, and workshops. It is used in Paris for lighting shops which require a light by which different colours may be distinguished, and has recently been used in England for the same purpose with satisfactory results. Many trials have been made for street illumination with greater or less success.

Compared with gas, the economy for equal illumination does not yet appear to be conclusively established. Although in some cases the relative economy for equal candle power is on the side of the electric light, yet in other cases gas illumination of equal intensity has the advantage. Unquestionably the electric light has not made that progress which would enable it in its present condition to enter into general competition with gas for the ordinary purposes of domestic supply. In large establishments the motors necessary to produce the electric light may be readily provided, but, so far as we have received evidence, no system of central origin and distribution suitable to houses of moderate size has hitherto been established.

In considering how far the Legislature should intervene in the present condition of electric lighting, the Committee would observe, generally, that in a system which is developing with remarkable rapidity it would be lamentable if there were any legislative restrictions calculated to interfere with that development. The Committee, however, are not in a position to make recommendations for conditions which may hereafter arise, but at present do not exist, as to the distribution of electric currents for lighting private houses from a central source of power. No legislative powers are required to enable large establishments, such as theatres, halls, or workshops, to generate electricity for their own use.

If corporations and other local authorities have not power under existing statutes to take up streets and lay wires for street lighting and other public uses of the electric light, your Committee think that ample power should be given them for this purpose. There seems to be some conflict of evidence as to whether the existing powers are sufficient or not. But even in regard to local authorities it would be necessary to impose restrictions upon placing the wires too near the telegraph wires used by the Post Office, as the transmitting power of the latter would be injuriously affected by the too close proximity of the powerful electric currents needed for producing light.

Gas companies, in the opinion of the Committee, have no special claim to be considered as the future distributors of electric light. They possess no monopoly of lighting public streets or private houses beyond that which is given to them by their power of laying pipes in streets. Electric light committed to their care might have a slow development. Besides, though gas companies are likely to benefit by the supply of gas to gas-engines which are well suited as machines for producing electric light, the general processes of gas manufacture and supply are quite unlike those needed for the production of electricity as a motor or illuminant.

The Committee, however, do not consider that the time has yet arrived to give general powers to private electric companies to break up the streets, unless by consent of the local authorities. It is, however, desirable that local authorities should have power to give facilities to companies or private individuals to conduct experiments. When the progress of invention brings a demand for facilities to transmit electricity as a source of power and light from a common centre for manufacturing and domestic purposes, then, no doubt, the public must receive compensation advantages for a monopoly of the use of the streets. As the time for this has not arrived, the Committee do not enter into this subject further in detail than to say that in such a case it might be expedient to give to the municipal authority a preference during a limited period to control the distribution and use of the electric light, and failing their acceptance of such a preference, that any monopoly given to a private company should be restricted to the short period required to remunerate them for the undertaking, with a reversionary right in the municipal authority to purchase the plant and machinery on easy terms. But at the present time the Committee do not consider that any further specific recommendation is necessary than that the local authorities

should have full powers to use the electric light for purposes of public illumination, and that the Legislature should show its willingness, when the demand arises, to give all reasonable powers for the full development of electricity as a source of power and light.

PROCEEDINGS OF SOCIETIES.

PHYSICAL SOCIETY.

Ordinary Meeting, June 14, 1879.

Prof. W. G. ADAMS, President, in the Chair.

New members:—Mr. Donald Macalister, B.A., and Mr. St. George Lane Fox.

Prof. MACLEOD described a plan for Suppressing the Induction Disturbances in a Telephone Circuit. It is known that a secondary battery, composed of metal plates and sulphuric acid, allows weak currents to pass while stopping those of high tension. Prof. Macleod therefore inserted a secondary battery of platinum plates between the line and the telephone, but this stopped both the induction and the vocal currents. When platinum wires were substituted for the plates, however, the induction currents were stopped, while the vocal currents could be feebly heard.

Dr. O. J. LODGE exhibited his New Reversing Key for Electrometer Work, which is preferable to the ordinary forms, as giving a high insulation, small capacity, and not requiring the hand to approach close to it to work it. It consists of four platinum wires, arranged in pairs crossing one another; one pair crossing between the other two. These are the terminals and contact pieces of the key. The middle pair are supported by an endless silk thread, which runs on two pulleys, one of which is fitted with a handle. On turning the handle to right or left the two middle wires are brought into contact with one or other of the two outer wires, and the current reversed at will. The whole is enclosed in a metal box.

Mr. J. F. MOULTON then demonstrated the results of the experiments of Mr. Spottiswoode and himself on the Sensitiveness of Electric Discharges in Vacuum Tubes. These experiments were undertaken to find the cause of the luminous layers or strata in the discharge, a Holtz machine being employed. It was observed that when feeble currents were drawn from the machine, the discharge could be depressed by laying the finger on the tube, and this depression always occurs with intermittent currents; therefore the feeble currents from a continuous current Holtz discharge themselves like intermittent currents by reason of their feebleness. This sensitiveness of the discharge to the approach of the finger was found to be due to the conductivity and electric capacity of the hand. Electricity, opposite in kind to the discharge, is induced on the finger, and streaming upon the tube neutralises part of the discharge therein. This effect was also shown by means of tinfoil rings round the tube. An intermittent current is of course capable of this static induction on neighbouring conductors. The luminous discharge in a vacuum tube consists of a bright sharp glow at the negative terminal, followed by a dark space, then a hazy bluish light at the positive pole. The striae or layers in these sensitive tubes merely repeat this appearance. They can be artificially produced by placing the fingers, or rings of metal, at intervals along a tube conveying an amorphous discharge; for in this case the induced electricity discharging itself from the fingers breaks up the amorphous discharge into dark and bright layers. In these stratified discharges the electricity appears to travel *per saltum*, or by stepping stones, as one may say, and the glow seems to be a molecular structure, a view which is supported by Mr. Crookes's

experiments. A negative discharge from the finger produces a dark space in the tube discharge, and a positive one a bright line; therefore, one can tell the kind of discharge passing in a tube by laying a finger on it. If the same pole be brought to both ends of a tube a discharge will still take place from each end, and there will be a dark space in the middle; the electricity here seeming to turn back again the way it came. The discharge from a pole through a vacuum tube would therefore appear to be not akin to conduction but to a disruptive discharge. It is a leap in the dark, and the phenomena observed are due to the gaseous nature of the medium. These experiments point to the possibility of completing a circuit by positive electricity alone.

Prof. GUTHRIE suggested that by combining vacuum tubes with the induction balance of Prof. Hughes it might be possible to get an optical balance for measuring inductive capacity.

Dr. HENRY DRAPER, of New York, who is now on a visit to England, then addressed the meeting on his alleged discovery of oxygen in the sun by bright lines in the solar spectrum. He said that hitherto he had not been able to find these lines projecting from the limb of the sun like hydrogen, and his impression is that oxygen resides lower than the reversing layer. He had lately been extending the dispersion of the spectrum of terrestrial oxygen, and from a light of maximum intensity of one candle power had now got a dispersion of 80 inches from A to O. He exhibited two of the original negatives of the solar spectrum, showing the bright lines.

Mr. J. NORMAN LOCKYER congratulated solar science on having so able a worker as Dr. Draper, and remarked that if Dr. Draper proved his case for even two or three O lines it would be sufficient, considering the variability of the spectrum of matter under different physical conditions. He also alluded to traces of carbon which he himself had found in the sun by the dark flutings in the spectrum.

Dr. DRAPER said he did not see why carbon should not give both bright and dark lines.

Mr. SCOTT exhibited a number of coloured photographs done after the method of Mr. Albert, of Munich.

NOTICES OF BOOKS.

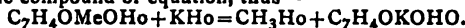
A Manual of Organic Chemistry. By HUGH CLEMENTS, of H.M. Civil Service, President of the Amateur Mechanics' Workshops Association, London, and Lecturer on Various Sciences at St. Thomas', Charterhouse, &c. Blackie and Son: London, Glasgow, Edinburgh, and Dublin, 1879.

THE difficulties of an examiner in setting questions, which shall fairly test the real knowledge of a candidate, have of late years considerably increased owing to the large number of text-books "adapted" to examinations which have been published, and serve merely to "cram" the candidate for the time being, with little or no regard to sound instruction. This is especially the case in respect to the examinations of the Science and Art Department. In a great measure, it is true, this is the fault of the system which makes the teacher dependent for his remuneration on the passing capabilities of his students, and therefore necessitates the use of those text-books which are best adapted for getting his men "through."

Such a book is the volume before us, forming one of a large class which has come into being with the rapid mushroom growth of the system of competitive examination. A feature in this Manual which cannot be too strongly condemned is the publication at the end of the book of a series of short answers to all the questions set in Organic Chemistry by the Examiners in the Science and Art Department during the last ten years. This

occupies no less than one-third of the whole volume. A student, therefore, who learns by rote a good selection of these answers runs a fair chance of passing, though his real knowledge may be of the most fragmentary and imperfect kind. As practical examinations are to a certain extent free from the objection which applies to written examinations, it is to be hoped that in the future much greater stress will be laid on practical work than has hitherto been the case. In the meantime it is highly satisfactory to know that the Department is alive to this necessity, and to find that a large percentage of the candidates in practical chemistry at these examinations send up very creditable results, showing that at many science schools a really sound knowledge of chemical analysis is being imparted.

As regards the book itself it is difficult to imagine how anyone who professes to be a teacher and to have experience enough to write a text-book on the subject could employ the arrangement adopted by the author. In many cases compounds are classed together which have little or no connection with one another; thus, under the heading of hydrides of organic radicals we have in the following order:—Hydrocarbons, cyanogen, oxalic acid, oxamic acid, oxamide, hydrocyanic acid, formic acid, and Prussian blue; whilst among organic bases we have cyanic acid. Nor is his plan of describing the apparatus employed in organic analysis, &c., to be recommended. This he does in a final chapter quite apart from the description of the processes themselves. Space also is here wasted unnecessarily in giving a figure of an ordinary "specimen tube" for weighing out the substance for analysis, as well as by drawings of two retorts, the difference between which it is impossible to see, except that in the one the neck is turned to the left and in the other to the right. The notation is exceedingly loose and defective, different symbols for the same or an analogous radical being frequently used indiscriminately, even in the same compound or equation, thus—



But one of the greatest defects in the book is the almost total neglect of the use of structural formulæ, without which a sound knowledge of organic chemistry can never be obtained by the ordinary student. In many even fundamental points the book is also very imperfect. General reactions are not made sufficiently prominent, and no explanation is given of isomerism and polymerism, nor of the difference between empirical and rational formulæ. To the glycols and their derivatives five lines only are devoted, whilst no special reference is made to glycerin, erythrite, mannite, &c., except as regards the identification of glycerin. The relations of the various organic acids to one another and to other compounds are by no means sufficiently insisted on, nor is any evident distinction made between primary, secondary, and tertiary alcohols, nor between the different behaviour of aldehyds and ketones on oxidation. The important series of aromatic compounds are passed over far too cursorily, no reference at all being made to the ortho-, para-, and meta-positions of the side chains in the derivatives of benzol, &c. Anthracen, naphthalen, and all the other important hydrocarbons and their derivatives, containing more than one benzol nucleus, are omitted altogether, except the identification of naphthalen.

But it is in the series of exercises given at the end of the book that the author shows his want of proper chemical training, or he would not set such childish questions as the following, in which and in many others he has evidently tried in how many different ways he could vary the same question:—"What organic substances are soluble in water?" "What insoluble in water?" "Which are soluble in alcohol?" "Which insoluble in alcohol?" "Which partly soluble in alcohol?" "Which insoluble in water but soluble in hot alcohol?" and so on, through quite a long string of such questions on solubilities. Then follows a long series of similar questions on the taste of organic substances, including—

"What organic substances taste peculiar?" More than one-third of all the exercises consist of such strings of questions, and would doubtless exercise the poor student to no small extent. Question (93) would be a puzzler in the way of analysis in most cases; it runs as follows:—"Make a qualitative analysis of the following mixtures:—Beef, or mutton, or blood, or milk, or flour, or oatmeal, or potatoes, or rice, or apples, or oranges, or tea, or coffee, or snuff, or opium, &c.

There are, however, one or two redeeming points, as the section on "Substitution," and especially that on "Organo-metallic Bodies," whilst the introduction of a special chapter on the identification of organic substances—a new feature in text-books on organic chemistry—has long been a desideratum.

CORRESPONDENCE.

ASHES OF WHEAT-BRAN.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxix., p. 244, there is an elaborate analysis of the ashes of wheat-bran. It seems to me that it is the "play of Hamlet with the character of Hamlet left out." In the analysis no mention is made of phosphate of alumina. As this compound plays so important a part in the analyses of flour and bread it ought not to have been left out. I have of late paid some attention to the analysis of bran from white English wheat, and I find that it contains—

Silica	3.671
Phosphate of alumina..	0.282
Phosphate of iron .. .	0.760

The percentage of sulphate of calcium should be 0.256, not 1.9567, as given in Mr. Peckham's analysis.—I am, &c.,

J. CARTER BELL.

VOLUMETRIC DETERMINATION OF CHROMIUM.

To the Editor of the Chemical News.

SIR,—In the last number of the Chemical Society's *Journal* there is the description of a process for determining chromium, based on the oxidation of chromium oxide by potassium permanganate in presence of sulphuric acid, also described in the CHEMICAL NEWS, vol. xxxix., p. 131. May I refer the author (Mr. Sell) to the CHEMICAL NEWS, vol. xxv., p. 151, in which I have described the same process for the determination of chromium in chrome iron and steel? The process as described there is in daily use, and gives thoroughly reliable results.—I am, &c.,

WM. GALBRAITH.

Atlas Works, Sheffield, June 10, 1879.

ON COLORIMETRY.

To the Editor of the Chemical News.

SIR,—At a meeting of the Literary and Philosophical Society of Manchester, on January 11, 1876, I proposed a method for estimating colouring matters in solution. Assuming that if we have in solution in equal bulks of a colourless menstruum the same colouring body, the lengths of the columns of fluid necessary to furnish the same colour are inversely as the quantities of the colouring matter present. Practically the method was carried out by the use of movable white disks in the cylinders containing the solutions. My experiments showed fair approximations when one liquid did not differ very much in strength from the other. Further experiments were made with fluids containing large quantities of colouring

matter, and with the white surfaces both external and internal, I find that the law holds through a wider range with the white surfaces external than internal. In a paper which I read before the Physical and Mathematical Section of the Literary and Philosophical Society in April of this year I pointed out the cause of this. As such a method of colorimetry is, I believe, in practical use, I think it well to bring the matter early under notice. The nature of the correction to be applied when the white surfaces are internal may be inferred from the following considerations:—The light which illuminates the surfaces has previously passed through the solution, so that really we are looking at a coloured disk through a coloured solution. Some allowance must be made for this colouration. If q and q' denote quantities of colouring matter, and t and t' the lengths of columns, then for external surfaces we may use the formula, $qt = q't'$. But suppose the surfaces are internal; then we must add to the lengths of the columns an additional length, which would produce the previous colouration. If the fluids compared do not differ much in composition an approximate formula is—

$$q(t+x) = q'(t'+x).$$

The value of x must be determined experimentally by comparing white surfaces seen externally and internally. If the fluids differ much in composition, the correction to be applied to the columns will differ a little on each side, so that a formula $q(t+x) = q'(t'+x)$ will be necessary. In the same paper I have pointed out a method of procedure in colorimetry which, I think, may be serviceable to those commencing experiments, or to those who are not good judges of colour, or in the case of colours which are difficult of comparison.

Given two cylinders containing coloured fluids of different intensities, and we wish to obtain the same tint in both. In the stronger solution start first with a tint darker than the other, and approach gradually to it; stop when the colours seem the same. Probably the columns will be a little too long. Next start in the stronger solution with a lighter tint than the standard solution; approach gradually to this limit, and stop when there seems no difference, probably the column will be too short. Next take the mean of the two columns.

The foregoing remarks apply to colouring matters in solution. A method of colorimetry founded on the same principle is, I believe, applied to turbid solutions. It is very desirable that before general adoption some exhaustive experiments should be made on the limits of its accuracy.—I am, &c.,

JAMES BOTTOMLEY, D.Sc.

Irwell Terrace, Lower Broughton,
near Manchester, June 16, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 21, June 2, 1879.

Magnetic Impenetrability of Iron.—J. Jamin.—A concentric thickness of 6 millimetres of iron arrests almost completely the magnetic effect of an external helix.

Ultra-violet Limit of the Solar Spectrum.—A. Cornu.—The utmost limit which the author has been able to reach corresponds to the wave-length 293.

Alkaline Amalgams and on the Nascent State.—M. Berthelot.—The author remarks that we possess hitherto very vague notions on the conditions of molecular mechanics which determine and regulate the use of alkaline amalgams. He has therefore undertaken their

thermic study. Sodium amalgam with water shows +32.6 cals., and potassium amalgam +27.0 cals. per equivalent of metal dissolved. With hydrochloric acid so diluted that 1 equiv. = 2 litres the results were for sodium +46.5, and for potassium +41.2 cals. The alkaline amalgams in hydrogenising reactions always disengage more heat than does free hydrogen. This thermic excess explains the hydrogenising power of amalgams, and throws a light upon the general theory of the reactions formerly ascribed to the nascent state.

Stanno-propyls and Iso-stanno-propyls.—MM. A. Cahours and E. Demarçay.—The use of an alloy containing less than 10 per cent of sodium yields a mixture of the diiodide of distanno-propyl and of the moniodide of tristanno-propyl, the separation of which is scarcely possible. To obtain the former of these compounds pure the product of the reaction of propyl-iodide upon tin must not be distilled except in a vacuum. Its composition is represented by $\text{Sn}_2(\text{C}_6\text{H}_7)_2\text{I}_2$. The iodide of tristanno-propyl is $\text{Sn}_2(\text{C}_6\text{H}_7)_3\text{I}$. The properties of these compounds are described at length.

Quantity of Nitric Acid Contained in the Water of the Nile before and after its Rise.—M. d'Abbadie.—The quantities of nitric acid per litre of water before, during, and after the floods are represented as being 0.01, 0.0038, and 0.002 grm.

Origin of Sound in the Telephone.—Th. du Moncel.—It is certain that the induction currents produced by a Bell's telephone can occasion the reproduction of speech in a magnetic disk surrounded with a helix without any need for a diaphragm influenced by such disk.

Rays of the Vapour of Sodium.—Mr. N. Lockyer.

New Arrangement for Increasing the Sensibility of the Vibrating Plate of the Telephone.—M. C. Decharme.—The author proposes to fix such plates by their centre only.

A Compound of Alumina and Carbonic Acid.—MM. Urbain and Renoul.—If alumina is precipitated from alum by means of sodic carbonate two bodies are obtained differing in aspect according as the liquid was boiling or of the common temperature of the atmosphere. In the former case the precipitate is gelatinous, transparent, and filters slowly, whilst in the latter case it is opaque, separates readily from the liquid, and absorbs colouring matters much more abundantly than the transparent variety. The author finds that the opaque variety contains carbonic acid in such proportions that it may be represented by the formula, $\text{CO}_2, 2\text{Al}_2\text{O}_3, 8\text{H}_2\text{O}$.

Mr. Lawrence Smith remarked that a natural carbonate of alumina and soda existed at Montreal.

Bulletin de la Société Chimique de Paris,
No. 10, May 20, 1879.

Oxy-ferrocyanide of Ammoniacal Copper.—Antony Guyard.—The precipitate obtained by adding potassic ferrocyanide to ammoniacal sulphate of copper if washed and dried in the air at about 150° loses cyanogen and ammonia, absorbs oxygen, and takes a violet tone. The material should be spread out in a thin layer, and stirred continually till the heat reaches 170°, when the reaction is complete. The colour does not seem available either in painting or in the pigment style in calico-printing. If heated up to 200° the colour changes to a blue, and at from 240° to 250° to a green, neither of them remarkable for brightness.

A Law Peculiar to the Metallic Ferrocyanides.—Antony Guyard.—The hydrated oxides, as existing in their soluble salts, may be divided into two great groups according to their behaviour with aqueous ammonia:—
1st. Oxides soluble in ammonia and in ammoniuretted ammoniacal salts, including, of course, the tartrate.
2nd. Oxides insoluble, or very sparingly soluble, in ammoniuretted ammoniacal salts, but, on the other hand,

soluble in ammoniacal tartrate. The former group is precipitated, and generally very completely, by potassic ferrocyanide, whilst those of the latter group remain perfectly dissolved in the basic ammoniac tartrate.

Volumetric-Chemical Studies.—W. Ostwald.—The contractions or dilatations experienced on mixing the solutions of salts may serve for the measurement of their respective affinities, and have furnished the author with results analogous to those obtained with the calorimeter by M. Berthelot and M. Thomsen.

Atti della R. Accademia dei Lincei.

Fasc. 4, March 1879.

Researches on Cinchonin.—Prof. Filetti.—The author dissolved 15 grms. cinchonin in a slight excess of hydrochloric acid, diluted the solution to 4 litres, saturated it with chlorine, and exposed it to the direct action of light. A white or slightly yellow substance was deposited on the sides of the beaker. This deposit dissolves in glacial acetic acid, and is re-precipitated by water. On heating cinchonin with bromine and water for several days in a closed tube he obtained products, the examination of which is still incomplete.

*Verhandlungen des Vereins zur Beförderung des
Gewerbflusses.* April, 1879.

This number contains no chemico-technological matter.

MISCELLANEOUS.

The Royal Society.—On Thursday, the 12th inst., the following fifteen candidates were elected Fellows of the Royal Society:—J. Anderson, M.D., Rev. M. J. Berkeley, H. Bessemer, Prof. A. Crum-Brown, W. L. Buller, Sc.D., G. H. Darwin, Prof. J. D. Everett, Prof. F. S. B. François de Chaumont, Prof. G. D. Liveing, G. Matthey, G. J. Romanes, A. Schuster, Ph.D., Prof. H. G. Seeley, B. Williamson, and T. Wright, M.D. The following have been elected Foreign Members of the Society:—Arthur Auwers, Berlin; Luigi Cremona, Rome; Jean Louis Armand de Quatrefages, Paris; Georg Hermann Quinke, Heidelberg; Theodor Schwann, Liege; Jean Servais Stas, Brussels.

NOTES AND QUERIES.

Liquid Perchloride of Iron.—I should be glad if any of your correspondents could tell me where there is a good market for liquid perchloride of iron in quantities (sp. gr. 1.400⁰), and the price that would be given. Also protochloride of iron (sp. gr. 1.400⁰). Have convenience for making large quantities.—SCARLET.

MEETINGS FOR THE WEEK.

MONDAY, 23rd.—Royal Geographical, 8.30.
TUESDAY, 24th.—Anthropological, 8.
WEDNESDAY, 25th.—Society of Arts, 4. (Anniversary.)
Geological, 8.
THURSDAY, 26th.—Royal Society Club, 6.30. (Anniversary.)
FRIDAY, 27th.—Quekett, 8.
SATURDAY, 28th.—Physical, 3.

THE YORKSHIRE COLLEGE.

The Worshipful Company of Clothworkers of the City of London having increased their Endowment of the Textile Industry Department of the College so as to include instruction in Dyeing, the Council of the College is prepared to receive applications for the post of Instructor; Stipend, £300 a year and half the Fees. Preference will be given to Candidates who are familiar with the processes of French and German Dyeing and with the Methods of Instruction pursued in the Continental Schools. Applications, accompanied by a statement of the Candidate's experience, together with copies of Testimonials, to be sent not later than August 11th to the Secretary of the Yorkshire College, from whom further information may be obtained. The work of instruction will begin in January, 1880.

Leeds, June 18th, 1879.

W. F. HUSBAND, Secretary.

Analytical Chemist desires an appointment.
Would be willing to make himself useful in any capacity; London preferred.—Address, Chemist, 351, Middleton Road, Oldham.

A German Chemist (Ph.D.), holding diplomas from University and Polytechnicum, who has already had experience in a manufactory, seeks a Situation in a Chemical Manufactory in England.—Address, K.D., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

Required by an Associate of the Royal School of Mines, F.C.S., &c., a Situation in a Laboratory, or as Manager of a Chemical or Metallurgical Works, or on a Mine, in England or abroad. Highest references.—Address, A.R.S.M., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

TO MANURE MANUFACTURERS.

FOR SALE.—About 150 tons Monthly of Woolen Shoddy; free from grease; containing nitrogen equal to from 5 to 10 per cent ammonia.—Apply to David Shaw and Co., Clayton, near Manchester.

Superior Iron Filter-Press for Sale, made to order and of extra quality, by Messrs. Needham and Kite. It contains ten chambers, each 19×21½ inches; is provided with a 3-inch gun-metal pump to work by hand or steam, and with fittings for washing and steaming, which can be used or not at discretion. The whole is quite equal to new, and is in perfect working order.—Address L. B. S., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

TANK WAGGONS.—Wanted two for Crocoite; secondhand if in good condition.—Full particulars to be addressed "Tank," CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

A Part or the Whole of Excellent Laboratory Fittings to be Sold with the Lease of an Eligible Dwelling-House. Rent very moderate.—Further particulars of E. and S. Smith, Auctioneers and Agents, 22, Southampton Buildings, Chancery Lane, W.C.

DAVID HILL, CONSULTING CHEMIST in Technical Processes, late of Deans Terrace, South Shields. Letters to be addressed care of Messrs. Bulloch and Co., 79, Mark Lane, London, E.C. Special attention given to questions relating to "Noxious Vapours."

WILLOUGHBY BROS., CENTRAL FOUNDRY, PLYMOUTH, Makers of Plant for Sulphuric Acid and Superphosphate Works, also for Tar and Ammonia Distilling.

NORRINGTON'S PATENT,

For the prevention of escape of gases during the charging of kilns and nitre pots. The success of this invention being assured, we are prepared to Contract for fixing this Patent Apparatus to Pyrites Furnaces throughout the Kingdom.

Estimates and Plans furnished on application.

Silicates of Soda and Potash in the state of Soluble glass, or in CONCENTRATED SOLUTION of first quality, suited for the manufacture of Soap and other purposes, supplied on best terms by W. GOSSAGE and Sons, Soap Works, Widnes, Lancashire.

London Agents, COSTE and Co., 19 and 20, Water Lane, Tower Street, E.C., who hold stock ready for delivery.

TO PROFESSORS & EXPERIMENTALISTS.

W. STONE, Mathematical, Surveying, and Optical Instrument Maker, 44, Gloucester Street, Queen's Square, Bloomsbury, works out all kinds of difficult experimental and scientific work.

S. A. SADLER, CLEVELAND CHEMICAL WORKS, MIDDLESBROUGH;

Newfall Tar Works, Carlton;
Ammonia Works, Stockton-on-Tees;
and Stamshaw Chemical Works, Portsmouth.

And also of the Furness Tar Products Co., Ulverston.

Manufacturer of Benzole, Toluole, Xylol, Solvent and Burning Naphthas, Carbolic Acid and Disinfecting Powder, Refined Anthracene, Naphthaline, Black Varnish, Refined Tar, Crude Liquid Ammonia, Coal-Tar, Pitch, Crocoite, Grease, Sulphate of Ammonia, Pyroligneous Acid, Acetate of Lime, Wood Naphtha, Charcoal, &c., &c.

S. A. S. is always a buyer of Coal-Tar Naphthas, Crude Anthracene and all Tar Products.

All communications to be addressed to the offices at Middlesbrough.

NOTICE—ARTIFICIAL ALIZARINE

London 18th June 1879

WE the undersigned the Owners respectively of the Patents for the Manufacture of Alizarine

A D 1869 No. 1936
A D 1869 No. 1948
A D 1869 No. 2218

HEREBY GIVE NOTICE That we recently commenced Legal Proceedings against The Alizarine and Anthracene Company Limited for damages for the infringement of the above-mentioned Patents and for an Injunction to restrain the said Alizarine and Anthracene Company Limited from manufacturing, selling or from offering for sale or disposing of or parting with any Artificial Alizarine or other Coloring Matters manufactured or prepared in contravention of our rights under the said Patents of either of them respectively and that after obtaining an interim injunction on the terms of our application we at the request of the said Alizarine and Anthracene Company Limited agreed to stay our Action upon the following terms

- That the said Alizarine and Anthracene Company Limited should submit to a perpetual Injunction
- That they should admit the validity of the said Patents
- That they should pay damages for past infringements
- That they should pay our Costs of the Legal Proceedings

A perpetual injunction against the Alizarine and Anthracene Company Limited have been granted and all proceedings in the Action have been stayed on the above terms

AND WE HEREBY GIVE FURTHER NOTICE that we recently were about to institute legal proceedings against Messrs. Arthur and Henshaw Drysalters and Oil Merchants in Glasgow and Matthew Clark and William Andrew Jamieson Drysalters and Oil Merchants Glasgow the individual partners of the said firm of Arthur and Henshaw to obtain interdict against their infringing the said Letters Patent or any of them and damages for infringements thereof by them in time past and that we have agreed to discontinue and discharge the said threatened proceedings upon certain conditions embodied in an Agreement entered into between us and them Under the said Agreement the said Messrs. Arthur and Henshaw and the said Matthew Clark and William Andrew Jamieson the individual partners of the said firm admit and acknowledge

- That the said Letters Patent are valid in every respect
- That they have infringed the said Letters Patent by importing and selling Artificial Alizarine manufactured according to one or other of the processes described in the Specifications respectively filed in connection with the said Letters Patent
- That the Letters Patent of the 28th May 1869 No. 1642 granted to Frank Julius Brönnner (therein named Julius Brönnner) and Hermann Gutzkow are and always have been null and void
- They undertake during the subsistence of our said Letters Patent not to import into or sell within the United Kingdom any Artificial Alizarine produced by any of the processes described in the Specifications of any of our said Letters Patent except with our consent in writing

WE LASTLY GIVE NOTICE that any person or persons infringing our above-mentioned Patents whether by purchasing (except from us or our Licensees) or selling importing or being concerned in importing or in anyway using Artificial Alizarine in the United Kingdom other than Alizarine made here by us or imported by us or our Licensees will be immediately proceeded against

A REWARD will be given to any person who will give the undersigned information sufficient to undertake legal proceedings against any such Infringers

BURT BOULTON and HAYWOOD
64 Cannon Street City London
Badische Anilin und Soda Fabrik
F. ENGELHORN AUGUST CLEMM

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Post free on application to JAMES K. GREGORY, Geologists and Mineralogists' Repository, 88, Charlotte Street, Fitzroy Square, London.

IMPORTANT TO SULPHURIC ACID MANUFACTURERS. NORRINGTON'S PATENT.

In the ordinary method of Manufacture, at the time of charging the Kilns, a considerable escape of gases takes place. This is attended with proportionate loss of Sulphur, and with much inconvenience to the workmen, as well as annoyance to the vicinity of the Works. This may be entirely avoided by the adoption of C. Norrington's patented invention, which can be applied at moderate cost to existing Plant, as well as in the erection of new Works. It may be seen in full operation on extensive Plant at Messrs. C. Norrington and Co.'s Chemical and Manure Works, Cattedown, Plymouth, where the fullest information may be obtained, with terms for cense.

THE CHEMICAL NEWS.

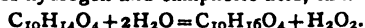
VOL. XXXIX. No. 1022.

LIMITED OXIDATION OF THE ESSENTIAL OILS.

THE ATMOSPHERIC OXIDATION OF TURPENTINE.

By E. T. KINGZETT.

1. In my previous papers,* which were communicated to the Chemical Society and the British Association, I have shown that when the so-called essential oils are exposed to atmospheric oxidation, peroxide of hydrogen is produced in an indirect manner. In the case of turpentine, which was more particularly studied, it seemed to me that a camphoric peroxide ($C_{10}H_{14}O_4$) is first formed, and that this is slowly decomposed by the action of water, yielding peroxide of hydrogen and camphoric acid, thus—



Certain it is that I obtained camphoric acid in my experiments, and analysed some of its compounds. I also succeeded in showing that all hydrocarbons of the formula $C_{10}H_{16}$ (terpenes) similarly give rise to the formation of peroxide of hydrogen, and that this property is shared by menthene, $C_{10}H_{18}$. On the other hand, it was observed that the hydrocarbons of the formula $C_{15}H_{24}$ did not behave in this way. The production of peroxide of hydrogen by the atmospheric oxidation of the specified hydrocarbons seemed to have some kind of connection with another property of the same bodies. It had been shown by Oppenheim and Wright that all the terpenes yield cymene ($C_{10}H_{14}$) when subjected to suitable processes, and, moreover, it appears that cymene as obtained from all sources is identical. According to Fittica, cymene is normal propyl-methyl-benzene, in which the methyl and propyl occupy the *para* position. This view has not received universal acceptance, but whatever it may be worth, there is apparently no satisfactory reason for doubting the identity of cymene prepared from different compounds, and the researches of Paterno lend particular confirmation to this statement.

Now, since all the terpenes and menthene give peroxide of hydrogen as above stated, and since they also contain a cymene nucleus, the two facts seem to be related, and this relation is confirmed by my observations, that cymene itself yields peroxide of hydrogen also, while the hydrocarbon from oil of cloves, $C_{15}H_{24}$ (Church), does not yield peroxide of hydrogen and (as I anticipated would be the case) does not yield cymene by bromination (as shown by Wright).

In the last part of my research the general features presented in the atmospheric oxidation of turpentine on a large scale were described in detail, and it was then shown that the compound which is primarily formed, and which afterwards (*viz.*, when in contact with water) gives rise to the formation of peroxide of hydrogen, may be obtained in such amount that when the oil of turpentine containing it is subjected to distillation decomposition occurs just above the boiling-point with almost explosive violence. As is well known, the atmospheric oxidation of turpentine is now conducted as a commercial operation (for producing the disinfectant called "Sanitas"), and I have been naturally anxious to continue my experiments in order to obtain a more perfect knowledge of the process and its products. Unfortunately, the performance of other duties has largely

interrupted my work, and even now I am only induced to make this preliminary communication, since, from a paper read at the last meeting of the Chemical Society, I perceive that Dr. Armstrong is working in some measure upon the same subject.

Having said so much, I will now give the notes of my recent experiments, all of which I intend to repeat and work out to their final conclusions as opportunity may offer and time permit.

2. There is a great difference presented by different essential oils, and even by various kinds of turpentine, as regards the rate at which they individually absorb oxygen from the air; and not only is this the case, but an oil of turpentine which, say, absorbs oxygen at a given rate, will absorb it much less rapidly if freed from contained oxidised matters by distillation. In other words, when once the process has begun, it continues more and increasingly rapidly, so that the greater the amount of oxidised matter contained in the oil the greater the oxidation under given conditions. The following table is interesting as illustrating the relative absorptive rates for atmospheric oxygen presented by the various oils experimented with. The method employed was as follows:—

A number of very long tubes (made from combustion tubing) and sealed at one end, were charged with equal volumes of air, water, and the various oils. After exposure to sunlight of summer for a number of hours, and before all the oxygen was absorbed in any case, the tubes were removed to the laboratory and the residual gas in each tube measured. Calling the largest amount of oxygen absorbed 100, then—

	Absorbed.
(a) Russian oil of turpentine	100.0
(b) Swedish " " " " " " " "	100.0
(c) Some oil which I obtained in Switzerland ..	89.4
(d) American oil of turpentine	78.9
(e) Oil of Eucalyptus	75.0
(f) An evidently adulterated Swedish turpentine	52.6
(g) " Scotch distilled American turpentine " "	42.1

The oils *a*, *b*, *d*, and *e* were undoubtedly genuine commercial oils; the oil *c* was perhaps French oil of turpentine, and in this case only was the oil distilled before use, because to my knowledge it had undergone great oxidation by keeping for many years; the oils *f* and *g* were evidently not genuine, and had been sent to me merely as samples; I should say that both had been adulterated with so-called pine oil of commerce. It may be mentioned that when the oils which absorbed atmospheric oxygen at the relative rates indicated by 100, 100, 78.9, and 75 were taken in equal amounts and placed in cylindrical glasses having their tops covered with paper treated with the same amount of the same mixture of potassic iodide and starch, the papers became coloured owing to the formation of peroxide of hydrogen in their vicinity in the same order. The same general result has been obtained in oxidising very large quantities of the same oils for commercial purposes.

3. In Parts II. and IV. I have described the general nature and characters of the aqueous solution which is obtained when oil of turpentine is, in the presence of water, exposed to the action of a blast of air, and it is therefore unnecessary to do the same thing again in this place. If this solution be evaporated on the water-bath to dryness, of course all the acetic acid is got rid of, and all the peroxide of hydrogen is decomposed. There remains behind a dark coloured matter, which when hot has a sugar-like odour and is viscid; on cooling it sets into an adhesive but firm mass, which, when treated with concentrated sulphuric acid, gives a colour reaction somewhat resembling that bearing Pettenkofer's name.

To get some sort of knowledge of its composition a part of the substance was dried at 100° C.,* and burnt with

* Part I.—*Journal of the Chemical Society*, June, 1874. Part II.—*Ibid.*, March, 1875. Part III.—*CHEMICAL NEWS*, vol. xxxii., p. 133. Part IV.—*Ibid.*, vol. xxxiv., p. 127.

* The weight did not become constant, the substance being evidently slowly volatile at 100° C. After being in the oven for forty or fifty hours the residual substance lost 10 per cent during the next forty hours.

romate of lead, using also in the tube a few pieces of stassic perchlorate so as to be sure of perfect combustion. 2450 grm. gave 0.2214 grm. H_2O and 0.5766 grm. CO_2 . These figures give—

	Per cent.	÷ at wts.	÷ 0 = 1.
C ..	64.18	5.3483	3.3
H ..	10.00	10.0000	6.1
O ..	25.83	1.6137	1.0

$\times 3 = C_{10}H_{18}O_3$

Of course this formula is only provisionally acceptable, but the information thus obtained is important.

This substance exercises remarkable antiseptic powers, and indeed the similar qualities of "Sanitas" are largely due to its presence, but they are supplemented by the properties of the accompanying peroxide of hydrogen and the substances which are dissipated during the evaporation of the solution.

When the substance whose analysis has been given above is heated with water it is found to be imperfectly soluble: roughly, about 5 per cent remains as a viscid, insoluble, nearly black matter. In one experiment, working with nearly half a pound of substance, this separation was effected, and for the time being the insoluble matter was set on one side. The yellowish brown solution was shaken with purified animal charcoal, but this failed to remove any of the colour, although the charcoal abstracted a considerable amount of the substance itself. The solution was finally evaporated to dryness, and there was thus obtained a transparent varnish-like substance, of an agreeable odour, and showing itself to be volatile when heated in the air-bath at $100^\circ C$. It was analysed, observing the same precautions as before, but the manipulation of the substance was very difficult, owing to its semi-fluid state when hot, and its adhesive nature in the cold.

0.2884 grm. gave 0.263 grm. H_2O and 0.6688 grm. CO_2 . These figures give—

	Per cent.	÷ at wts.	÷ 0 = 1.
C ..	63.245	5.2704	3.17
H ..	10.132	10.132	6.09
O ..	26.623	1.663	1.00

$\times 3 = C_{10}H_{18}O_3$

I believe that in this analysis the carbon came out a little too low on account of the difficulty just mentioned, but practically it may be said that the removal of more than 5 per cent of matter from the originally analysed substance did not materially affect its composition. It will be convenient here to speak of that part of the residue soluble in water as the soluble part, and that which is left undissolved by water as the insoluble part.

The insoluble part does not give the vivid reaction with strong sulphuric acid that is afforded by the other part, and even when a drop of strong sugar solution is added but little purple colour is obtained. Although insoluble in water it readily dissolves in an oily matter (to be hereafterwards described) which accompanies it and the other substance in the original solution; hence its insolubility is explained.

The soluble part, $C_{10}H_{18}O_3$ (?), gives with strong sulphuric acid a deep cherry-red colour reaction, and this is not intensified by adding syrup. When heated on platinum-foil it melts, evolves inflammable gas, and leaves a charcoal. Its aqueous solution reduces Fehling's solution just as glucose does, and exhibits very powerful antiseptic characters.

4. A quantity of this substance was subjected to distillation. At first it melted, then boiled, and a small quantity of an almost colourless oil passed over, which condensed into a colourless, soft, crystalline mass on cooling. After this, permanent oil passed over, which grew denser and darker as the distillation proceeded. The vapours in the retort seemed at the end to have a green colour. When the distillation was stopped there remained in the retort a black-looking liquid mass, which set on cooling, and then resembled pitch in appearance. The experiment was quite a preliminary one, and I have not yet repeated the

process more carefully with a view to determine the temperatures at which the different products came over; neither have I yet examined, analytically or otherwise, either the crystalline matter or the liquid oils.

Again, if the body whose analytical numbers agree with the formula $C_{10}H_{18}O_3$ be dissolved in water, and the solution acidified with dilute sulphuric acid, it grows milky, and on standing a slightly coloured oily body of considerable quantity separates. This reaction will probably lead on further examination to a better understanding of the constitution of the substance $C_{10}H_{18}O_3$; and indeed I have no doubt that with a knowledge of this and the products of its change there will be obtained very important evidence as regards the nature and constitution of the terpenes as a class.

5. Several litres of the aqueous solution ("Sanitas") obtained by oxidising Russian turpentine in the presence of water, were treated with caustic soda in bare excess and the mixture, which darkened very much at the point of neutralisation, was evaporated nearly to dryness on the water-bath. The dark-coloured soft resin-like residue thus obtained was then treated with an excess of dilute sulphuric acid. By this treatment there separated a dark oily mass, from which the clear solution was filtered, and then subjected to distillation. As the acid solution became hot more oil separated out, and there passed over an acid distillate accompanied by 20 or 30 c.c. of a slightly yellow volatile oil, with an odour resembling that of mircymene and eucalyptus. This oil, which gives a beautiful colour-reaction with strong sulphuric acid, is probably identical with that obtained in (4) by treatment of $C_{10}H_{18}O_3$ substance with dilute sulphuric acid. At the end of the distillation there remained in the retort a quantity of black tarry matter floating in a fluid state on surface, and on cooling it set into a thick skin. The acid distillate was separated from the oil, and then exactly neutralised with pure caustic soda, and the solution evaporated to the point of crystallisation. In this way crystals of sodic acetate were obtained, and subsequently pure acetic acid and its other salts were made. The acetic acid obtained did not amount to more than 0.25 grm. per litre of fluid operated upon; nevertheless, this observation confirms similar facts which I stated in the second part of this research. I could detect no other volatile acid.

If the aqueous solution which formed the subject of this last experiment be distilled alone, a certain amount of oil also passes over with the distillate. For instance, in one experiment 1000 c.c. fluid yielded about 5 c.c. of oil. More than double the amount, however, is obtained if the solution be acidified with sulphuric acid. Whether, however, these oily distillates are identical in composition cannot yet be stated, nor am I as yet assured that no other volatile matter soluble in water also passes over.

6. I have now sufficiently indicated the work that has lately occupied my attention, and I may add that I anticipate its further prosecution will be attended with very interesting and important results. This is particularly the case, since in my process of oxidation there is no violent change like that resulting from the use, for example, of nitric acid, and hence the products are much more nearly related to and comparable with the original terpene employed: as a consequence the results are more intelligible and the inferences to be drawn from them acceptable accordingly.

7. It is much to be regretted that the recorded information regarding other processes of the oxidation of turpentine and their products is not more complete and precise, for it is mainly by the light of such knowledge that chemists trust to ascertain the constitution of this and the allied hydrocarbons. I trust I may be excused for saying chemists as a body are too fond of confining their studies to a limited number of the products of any one process, and they preferably choose of course the most crystalline of these. This is unfortunate, because any one reaction which may be worked out is not rarely regarded as the only one which occurs in a given process, whereas the

truth is more often that it is only one of a large number that really occur.

When turpentine is oxidised by heating it with plumbic oxide, it is said to yield terebinic and formic acids, thus:—
 $2C_{10}H_{16} + 7O_2 = 2C_9H_{14}O_5 + 2CH_2O_2$.

This reaction, undoubtedly, does not express the whole truth. Then, again, Hempel has stated that when turpentine is oxidised with chromic acid it yields terpenylic acid, $C_9H_{12}O_4$. This is an important observation, but would be much more valuable if all the accompanying products were also studied and defined.

By oxidising turpentine with nitric acid, toluic acid, $C_8H_8O_2$, and terephthalic acid, $C_8H_6O_4$, are also said to result, but Wright has raised the question whether the terephthalic acid thus obtained is not due merely to the presence of cymene in the turpentine operated upon.

More recently, however, Hempel has repeated the statement that these acids are obtained by the process above indicated. They are also said to be accompanied with terebic acid, $C_7H_{10}O_4$, terebenzic acid, $C_7H_7O_2$, and terechrysic acid, $C_6H_8O_5$.

This variety of products attending the use of such powerful oxidants as nitric acid or an acid solution of potassic dichromate seems clearly to establish the great advantage of employing the mildest possible oxidising agent where it is desired to obtain a knowledge of the constitution of substances. Indeed, the presumption is that, since it is possible to resolve turpentine into carbonic anhydride and water, if the oxidation with nitric acid be continued for a sufficient period, not only the products previously indicated but a further vast number may be intermediately produced.

ON SOME ANALYSES OF IRON.

By SERGIUS KERN, M.E., St. Petersburg.

In several articles inserted in this journal the author has pointed out that in many cases the chemical analysis of iron or steel cannot give an idea of the quality of the metal. The following is a short account of some experiments executed some time ago.

Some pieces of rolled iron (boiler plates) prepared from a very pure pig-iron were sent for analysis. The iron, however, proved to be of a bad quality, and could not stand all the necessary tests. The following are the results of the analysis:—

Silicon	0.010 per cent
Manganese	0.120 "
Sulphur	none "
Phosphorus	trace "
Copper	0.028 "

This analysis cannot show that the inferior quality of the iron was due to the presence of noxious elements. The author believes that the small quantity of copper could not have any influence on the quality of the iron. The production of bad plates from such a beautiful iron could take place only supposing that the rolling of the metal was badly conducted.

ON NITRIC NITROGEN IN MANURE.

By BEAUMONT J. GROSJEAN,
J. B. Lawes's Chemical Works, Millwall, E.

MR. TATLOCK (p. 266) has evidently not seen some results which I published in the *CHEMICAL NEWS*, vol. xxv., p. 205, on the above subject, since he states that he is not aware that the complete decomposition of the nitrates in guano—in the course of determining the nitrogen present—has till now been accomplished, or even the necessity for it admitted. My experiments were simply preliminary.

I did not follow them up, owing to the fact that about that time manures ceased to be analysed in our laboratory. I showed that the *partial* decomposition of nitrates by soda-lime was well known, and was stated in, *e.g.*, Fresenius's "Quantitative Analysis" and Church's "Laboratory Guide." I described encouraging results obtained by making a combustion of nitre with iron filings and sugar. I was more successful when I adopted the following plan:—The nitre was dissolved in caustic alkaline solution, iron filings added, and distillation of the mixture to a pasty state carried on in a retort. The results, in fact, erred on the side of excess. I proposed that, in a mixture containing nitrogen in the form of ammonia salt, nitrate, and organic matter, the distillation should be carried on to the above point, and that then the pasty residue should be cooled. In this state it solidifies, and may be powdered, mixed with soda-lime, and a combustion of the mixture made to determine the residue of the organic nitrogen. I should be glad if Mr. Tatlock would carry out this idea.

RELATIONS BETWEEN THE ATOMIC WEIGHTS AND CERTAIN PHYSICAL PROPERTIES

(MELTING- AND BOILING-POINTS AND HEATS OF FORMATION
OF ELEMENTS AND COMPOUNDS.*

By THOMAS CARNELLEY, D.Sc.,
Assistant Lecturer on Chemistry in Owens College, Manchester.

THE object of the present paper is to trace the influence of the atomic weights on the melting- and boiling-points and heats of formation of elements, and especially of their compounds. It is shown that, as regards the elements, the melting-points are a periodic function of their atomic weights, whilst, for compounds, the following conclusions are drawn:—

I. That the melting- and boiling-points and heats of formation of the normal halogen compounds of the elements are a periodic function of the atomic weights of the constituent elements.

II. That the influence of the halogen on these same physical properties increases with the number of its atoms in the compound.

III. That in any normal halogen compound the influence of either of the elements on the melting- or boiling-point increases with its own atomic weight, and decreases with the atomic weight of the other element.

IV. The melting- or boiling-point or heat of formation of a bromide is always nearer to that of the corresponding chloride than to that of the corresponding iodide; and that the melting- or boiling-points of the halogen compounds of the middle member of three consecutive elements of the same group are always nearer to those of the first member (*i.e.*, the one with least atomic weight) than to those of the last member.

The former of these phenomena probably depends on the fact that the atomic weight of Br is nearer to that of Cl than to that of I, *i.e.*, less than the mean of the two; and the latter on the fact that the atomic weight of the middle member of three consecutive elements of the same group is always less than the mean of those of the other two elements.

V. That the melting- and boiling-points of the halogen compounds of the elements belonging to the first and second groups of Mendelejeff's classification are widely separated from those of the other groups, being in fact considerably higher. Different relations too often appear to exist between the melting-points of the *even* members of these two groups to those which exist between groups (3 to 7) inclusive; while the compounds of the elements, which are often placed in the *odd* divisions of the first and second groups, are generally altogether irregular. In

* Abstract of a Paper read before the Royal Society, June 17

the case of the odd members of the first group this may be explained by the fact that it is very uncertain whether Cu, Ag, and Au really belong to the same group as Na or not, as pointed out by Mendeleeff in his memoir on the "Periodic Law," in which he places these elements not in the first group along with Na, but in the eighth with Fe, Pd, Pt, &c.

As far as existing data allow us to judge, the compounds of the elements with monatomic organic radicals also obey the same laws as those of the halogen compounds.

A process for calculating unknown melting- and boiling-points (in the case of halogen compounds) by a method of limits is next described, which gives very good results when applied to known melting- and boiling-points, the average error for all known boiling-points being $\pm 4.5^\circ \text{C.}$, or 0.8 per cent, and for all known melting-points ± 15 , or 3 per cent. The calculated values for a large number of unknown melting- and boiling-points are given.

It is also shown that a knowledge of the melting- or boiling-points of the halogen compounds of an element may serve for the determination of the atomic weight when the application of the methods of specific heat and vapour-density are inadmissible, or the results obtained by these contradictory.

An application of this is made in the case of beryllium, concerning the atomic weight of which there has recently been some dispute. From his determination of the specific heat of the metal, Emerson Reynolds concludes that it is a dyad with atomic weight 9.2, whereas Nilson and Petterssen, from their determination of the specific heat, assign to it the atomic weight 13.8, in which case it would be trivalent. Now, according to calculation, the melting-points of the chloride, bromide, and iodide of beryllium ought to be $(820^\circ \text{ to } 870^\circ)^*$, $(802^\circ \text{ to } 820^\circ)$, and $(766^\circ \text{ to } 770^\circ)$ respectively. The boiling-points, also, ought to diminish from the chloride to the iodide, if beryllium is a dyad with atomic weight 9.2. If it be a triad with atomic weight 13.8 these melting-points ought to be about 500° lower, and the melting- and boiling-points ought to increase from the chloride to the iodide. With the object of settling this point the melting-points of the chloride and bromide have been carefully determined, and it was found that the chloride melts at 858° to 890° , thus agreeing with the limits $(820^\circ \text{ to } 870^\circ)$ calculated for BeCl_2 ($\text{Be} = 9.2$). The bromide also fuses between 858° and 890° , and at almost exactly the same temperature as the chloride, but if anything slightly higher, the calculated number for BeBr_2 being 802° to 820° . The rather high number found for the bromide is probably due to the substance being so readily volatile below its melting-point that the heat absorbed during conversion into vapour cools the remainder of the solid, and thus prevents it melting so soon as it otherwise would do. These results confirm the view that beryllium is a dyad, with atomic weight 9.2; for, though the melting-point of the bromide is apparently slightly higher than that of the chloride, for the reason already given, yet BeBr_2 is far more volatile than BeCl_2 .

ON THE PRODUCTION OF COLOURED SPECTRA BY LIGHT.†

By Captain ABNEY, R.E., F.R.S.

LAST year I incidentally mentioned in a note to the Royal Society (*Proceedings*, vol. xxvii., p. 291) that the production of natural colours by the agency of light, examples of which were shown by Becquerel, was probably caused by the oxidation of silver compounds employed. I have ventured to return to the subject in order to show

that the colours are so produced and are not due to interference.

I have sent, for the Society's inspection, pictures of the solar spectrum on silver plates, and also on compounds of silver held *in situ* by collodion. It will be observed that the spectrum has imprinted itself in approximately its natural colours; that on the silver plates it is more brilliant than on the collodion film, but that in the latter the colours are seen by transmitted as well as by reflected light.

I reserve for the present the exact details of the production of these pictures, but may say that they are produced by oxidation of silver compounds when placed in the spectrum; an exposure of two minutes being amply sufficient with a wide slit to impress the colours. The colouring matter seems to be due to a mixture of two different sizes of molecules of the same chemical composition, one of which absorbs at the blue end and the other at the red end of the spectrum, and the sizes of the molecules are unalterable whilst exposed to the same wave-lengths as those by which they were produced. I believe it possible and probable that the colours may be preserved unchanged when exposed to ordinary daylight.

ON THE FORMATION OF HYDROCYANIC ACID IN THE ELECTRIC ARC.*

By JAMES DEWAR, M.A., F.R.S., Professor of Chemistry to the Royal Institution.

A SERIES of experiments favouring the conclusion that the so-called carbon lines are invariably associated with the formation of acetylene† induced me to make some experiments to ascertain whether this substance can be extracted from the electric arc, which invariably shows this peculiar spectrum at the positive pole, when it is powerful and occasionally intermittent. For this purpose the carbons were used in the form of tubes, so that a current of air could be drawn by means of an aspirator through either pole, and the products thus extracted from the arc collected in water, alkalies, and other absorbents. Gases may be led through one of the poles, and suction induced through the other, in order to examine their effect on the arc.

The following experiments record the results obtained by means of the Siemens and de Méritens' magneto-machines:—

Experiment 1.—Drew a current of air by an aspirator through the drilled negative carbon, and passed the gases through potash, and iodide of potassium, and starch paste; found no nitrites; potash contained sulphides.

Experiment 2.—Hydrogen led in by the positive pole and the gases extracted as above, gave the well-known acetylene compound with ammoniacal sub-chloride of copper; while at the same time, a wash-bottle containing water gave distinct evidences of the presence of hydrocyanic acid.

Experiment 3.—Hydrogen flame burning alone gave no sulphides or hydrocyanic acid, the condensed water in a small bulb gave nitrites.

Experiment 4.—Air drawn through the negative carbon gave considerable quantities of hydrocyanic acid, which was greatly increased by extracting the gases through the positive carbon. Air was aspirated at the rate of about one litre per minute.

Experiment 5.—The same carbons used with de Méritens' magneto-machine gave no result.

Experiment 6.—Carbons purified in chlorine and hydrogen gave with de Méritens' nothing; with Siemens' a draught of air through the negative pole, a small quantity of hydrocyanic acid, but a larger yield when

* Reckoned from absolute zero -273 .

† A Paper read before the Royal Society, June 19, 1879.

* A Paper read before the Royal Society, June 19, 1879.

† As suggested by Plucker, Angstrom, and Thalén.

positive used. The gases extracted after the absorption of the hydrocyanic acid contained acetylene. If the carbons are not purified, sulphuretted hydrogen is always found along with other gases.

The inference to be drawn from the above experiments is that the high temperature of the positive pole is required to produce the reaction, which is in all probability the result of acetylene reacting with free nitrogen, as when induction sparks are passed through the mixed gases, viz., $C_2H_2 + N_2 = 2HCN$, and that the hydrogen is obtained from the decomposition of aqueous vapour, and the combined hydrogen in the carbons. It is possible, traces of alkaline salts in the carbon poles may favour the formation of hydrocyanic acid, but, as all attempts to purify the poles so as to stop the reaction failed, I am inclined to believe it is a direct synthesis. The acetylene reaction is one of the many remarkable syntheses discovered by Prof. Berthelot of Paris. The presence of sulphuretted hydrogen is doubtless due to the reduction of the sulphates, invariably present in the ash of the carbon.

A more complete examination of the various reactions to be brought about by means of the Siemens arc, and with poles of varied composition, and in presence of different gases, will be communicated to the Society in a subsequent paper.

My thanks are especially due to Messrs. Ausdell and Cottrell, assistants in the Royal Institution, for their valuable aid.

ON THE SMOKE OF AN ELECTRIC LAMP.*

By B. S. PROCTOR.

At our meeting in December, 1878, Mr. J. W. Swan exhibited an electric lamp, on the incandescence principle, which had broken down in consequence of the electric force being too great for the cylinder of carbon through which it had to pass. A note on this subject appears in the current part of our *Transactions*, p. 190. One of the points of interest noted was the appearance of a sooty deposit on the inside of the glass. The flask which contained the carbon pencil and its platinum conductors, having been filled with nitrogen and exhausted with a Sprengel pump, was supposed to contain nothing which could act as a carrier to convey by chemical means any carbon from the incandescent pencil to the cooler surfaces in its neighbourhood. The phenomenon appeared to be such as has been spoken of under the term "volatilisation of carbon." Mr. Swan having placed the lamp at my disposal for examination, I have now the pleasure of bringing under your notice the results.

Under the microscope the smoky deposit on the glass showed numerous bright globules, no doubt platinum, and more minute particles of dark matter nebulous under a $\frac{1}{2}$ -inch objective.

A fragment of the glass exposed to an oxidising heat, the deposit partially disappeared, still leaving the glass slightly darkened.

The platinum support—which had also a coating of dark sublimate at a little distance above and below the carbon pencil, but not in immediate juxtaposition with it—was next examined by exposing to the blowpipe flame the unsmoked portion, so that the conducted heat might act upon the deposit without the fear of the blast carrying away the matter, which was very loosely attached. In this way the deposit was burnt off without the mechanical action of the blast, the heat to which it had been subjected being that of dull redness.

A piece of the glass was then treated with aqua regia for several days. The deposit was diminished, but far from being entirely dissolved; the solution gave a blue reaction with yellow prussiate of potash, and no coloura-

tion with tannin till aided by vapour of carbonate of ammonia, when the usual purple colour of ferric tannate was developed. There is thus evidence of the deposit containing platinum, carbon, and iron. Probably the scattering of platinum globules might result from the disruptive discharge which took place at the moment of the lamp breaking down.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, June 19, 1879.

Dr. Roscoe, Vice-President, in the Chair.

THE minutes of the last meeting were read and confirmed. The following certificates were read for the first time:—J. W. Smith, W. R. Eaton Hodgkins, J. R. Ashwell, J. Bemrose. During the evening the following gentlemen were ballotted for and declared by the scrutators, Messrs. Carteighe and Tribe, to be duly elected Fellows of the Society:—C. J. Wilson, R. Reid, G. R. Tweedie, W. T. Bayne, J. Fletcher, H. Appleby, J. Sakurai, A. E. Robinson. The following papers were read:—

"On Gardenin," by J. STENHOUSE and C. E. GROVES. A short note on this substance was published in the Society's *Journal* some time ago by the authors, in which they described the method of extracting this substance from the so-called "Dekamali gum," which is the resinous exudation from *Gardenia lucida*; they also obtained a red compound by the action of nitric acid. In the present paper the authors continue the above investigation. The resin of the *Gardenia lucida* has a peculiar and unpleasant alliaceous odour, which is undoubtedly due to some volatile compound. By distilling the resin in a current of steam a volatile oil came over. The bulk of this distilled at 170° , and, after rectification over sodium, yielding a terpene boiling at 160° , having the formula $C_{10}H_{16}$; a small quantity of the oil boiled at 250° . The residue consisted of a dark brown liquid, having an aromatic and slightly alliaceous odour, and containing a trace of sulphur. Details of the methods employed for extracting and purifying the gardenin are given in the paper. The pure substance has the formula $C_{14}H_{12}O_6$. On treating 1 part of finely divided gardenin with 10 of nitric acid, sp. gr. 1.24, keeping the mixture cool, the whole turns of an orange colour, and finally solidifies in a few minutes to a paste of bright red needles, which are collected, washed rapidly to prevent further decomposition, dried, and crystallised from boiling chloroform; this substance has been provisionally named gardenic acid, though the authors have no evidence that it is really an organic acid, its behaviour with reducing agents pointing rather to a quinon grouping. It is insoluble in water, light petroleum, and carbon bisulphide, and almost insoluble in ether and benzene; dissolves readily in cold dilute alkaline solutions with a deep yellow colour, being re-precipitated on the addition of acids; its formula is $C_{14}H_{10}O_6$; it melts with decomposition at 223° . When a small quantity of strong nitric acid is added to a cold supersaturated solution of gardenin in glacial acetic acid, a mass of crystals of gardenic acid is obtained. If, however, this product, after being washed and dried, is boiled for some time with glacial acetic acid and re-crystallised from that solvent, it becomes more orange-coloured and melts at 243° , and dissolves in alkaline solutions but sparingly in the cold, yielding a purple solution, turning pale red on boiling. This acetyl compound dissolves readily in boiling alkaline solutions, from which it is precipitated by acids; it has the formula $C_{14}H_8O_6(C_2H_3O)_2$. When gardenin is left in contact with dilute nitric acid, sp. gr. 1.24, for a con-

* A Paper read before the Newcastle-upon-Tyne Chemical Society, March 27, 1879.

siderable time, gas is evolved, and a bright orange-coloured substance is formed, which is seen by the microscope to consist of colourless crystals mixed with an orange amorphous substance. The crystals are soluble, the amorphous substance insoluble in water. In contact with an aqueous solution of sulphurous acid, gardenic acid acquires a bright yellow colour and is converted into hydrogardenic acid, $C_{14}H_{14}O_6$; it forms flat lustrous needles resembling lead iodide; it melts at 190° , and is insoluble in water, light petroleum, and carbon disulphide, sparingly soluble in ether, easily soluble in hot benzene, spirit, or glacial acetic acid; it dissolves in dilute alkaline solutions, from which it is precipitated by acids; oxidising agents reconvert it into gardenic acid. Further researches as to the constitution of gardenic acid and gardenin are promised.

"On Dry Copper-zinc Couples and Analogous Agents," by J. H. GLADSTONE and A. TRIBE. The authors, in order to avoid the time, trouble, and the use of alcohol and ether necessary for the formation of the ordinary zinc couple prepared by immersing zinc foil in a 2 per cent solution of copper sulphate, &c., have turned their attention to the production of a dry couple as active as the moist couple. After numerous experiments the following method was found to be most satisfactory:—9 parts by weight of coarse zinc filings are placed in a dry flask with 1 part of finely divided copper; the mouth of the flask is then closed by a cork, through which passes a tube having a capillary termination. The metals are mixed and heated in a Bunsen flame with continuous shaking until the filings begin to lose their shape and acquire a yellowish tinge; the mixture is then somewhat rapidly shaken first in and then out of the flame. If the operation has been successful a number of small dark gray granular masses are obtained. The state of division of the copper is of great importance; the authors recommend the copper obtained by reducing the protoxide at the lowest possible temperature in hydrogen and sifting the product through fine muslin. 10 grms. of a dry couple prepared as above convert 5 c.c. of ethyl-iodide into zinc ethiodide in about six minutes when heated to 90° C. The authors in one experiment employed 87 grms. of ethyl-iodide, with a proportionate quantity of couple; in fifteen minutes a conversion equal to 90.4 per cent of the theoretical amount had taken place. The authors have, in the second portion of the paper, investigated the action of other couples. Zinc and platinum form a couple which decomposes water more energetically than the copper-zinc couple. In the preparation of dry couples no combination was found to be superior to the copper-zinc combination, probably because the copper sticks better to the zinc than either platinum, silver, or gold; it is also less liable to form alloys with zinc than these three metals. Magnesium and platinum form an exceedingly active wet couple, but an effective dry couple could not be obtained. Zinc with either cuprous or cupric oxide forms a dry couple almost equal in activity to that formed by zinc and metallic copper. The authors conclude that for practical purposes no combination is superior to that formed with zinc and copper prepared as above described.

In answer to Mr. GROVES,

Dr. GLADSTONE did not think that any difficulty would be found in preparing zinc-ethyl on a manufacturing scale with the aid of the copper-zinc couple.

Mr. WANKLYN had been much struck with the statement of the authors that when the copper and zinc were alloyed the couple so formed was inactive; it seemed to him that some slight alteration of the surface of the zinc was at the bottom of these reactions. Some long time ago he had left some ethyl-iodide in contact with ordinary zinc dust for some time in sealed tubes at the ordinary temperature: on opening them the conversion was found to be almost complete. The late Mr. Chapman and himself had observed that magnesium amalgam decomposed water, whereas sodium amalgam was comparatively inactive.

Dr. GLADSTONE suggested that the cause of the de-

composition referred to by Mr. Wanklyn was the metallic impurities present in ordinary zinc dust, lead, &c.

Dr. ARMSTRONG then read a paper by Dr. TILDEN and himself on "*The Action of Sulphuric Acid on the Hydrocarbons of the Formula $C_{10}H_{16}$* ." The authors describe many experiments on the action of sulphuric acid of various strengths and at various temperatures, chiefly on the terpenes of American and French turpentines. The so-called terebene can be best prepared by using concentrated acid at about 80° to 90° ; at lower temperatures much polymerisation takes place. The distillate obtained by passing steam through the product of the above action is not, as Ribau has stated, a mixture of cymene with an optically inactive liquid isomeride of terpene; the so-called terebene being really inactive camphene. By long fractional distillation it is obtained as a crystalline solid, melting after crystallisation from alcohol at 47° . The distillate contains also much of a $C_{10}H_{16}$ hydrocarbon, boiling at 176° , which is terpinene. The product of the action of dilute sulphuric acid on terpene is terpinene without camphene. The crude colophene which is not carried over by steam yields on distillation 10 to 30 per cent of substances volatile in steam, principally inactive camphene and terpinene, also cymene, a small quantity of a paraffin-like body, an optically inactive camphol (borneol), $C_{10}H_{17}OH$, &c.

"Researches on the Terpenes, Camphor, and Allied Compounds" (Part I.), by Dr. ARMSTRONG. On Hydrocarbons associated with the Terpenes, and on the Formation of Cymene from Terpenes and Allied Compounds.—A sample of terebene, given to the author by Dr. Hugo Müller, consisted almost entirely of cymene and of a paraffin-like body, boiling at 170° , but having the composition $C_{10}H_{20}$. On treating various terpenes with sulphuric acid similar results were obtained, in most cases 3.5 per cent of cymene and 0.5 per cent of the hydrocarbon insoluble in concentrated sulphuric acid. Russian turpentine contains 8 per cent cymene and 2 per cent of the hydrocarbon. Terpinene, resin spirit, the distillate from india-rubber, that from colophene, contain much cymene, with this insoluble (in sulphuric acid) hydrocarbon. The author believes the reaction $3C_{10}H_{16} = 2C_{10}H_{14} + C_{10}H_{20}$ takes place under these circumstances.

Part II. On the Action of Iodine on Terpenes.—The product obtained by distilling turpentine with one-fourth its weight of iodine is not merely cymene or so complex as the product of Preis and Raymann (*Ber. der Deutsch.*, xii., 219). It contains 33 per cent of hydrocarbons insoluble in sulphuric acid, apparently a mixture of two hydrides of the formula $C_{10}H_{20}$, boiling at 160° and at 170° . The portion soluble in acid contains no appreciable amount of lower homologues of cymene, but consists of camphene, cymene, and higher homologues. A small quantity of what appears to be a mixture of methyl and ethyl iodides is also produced in the reaction.

Part III. "*Camphor Derivatives*," by Dr. ARMSTRONG and Mr. MATTHEWS. The body previously described, as formed with camphoric acid by heating bromo-camphor with nitric acid, is a nitro-derivative. Dibromo-camphor yields no camphoric acid on oxidation with nitric acid, but is slowly and completely decomposed; alcoholic potash reduces it to mono-brom-camphor. Chlorine at 100° converts camphor into mono- and dichlor-camphor. The former resembles bromo-camphor in its behaviour with nitric acid; heated with alcoholic potash it retains its chlorine, and exchanges its hydrogen for hydroxyl. With bromine, chloro-camphor yields a magnificently crystalline bromo-chlor-camphor; these bodies resist oxidation. The above nitro-derivatives with alcoholic potash give potassic nitrite and hydroxy derivatives. Iodo-camphor is readily prepared by the action of iodine chloride on camphor. Mixed iodo-derivatives are also obtained in the same way.

After a few remarks by Dr. Wright, Mr. H. F. BROWN read an abstract of a lengthy paper by Mr. HERON and himself, entitled "*Contributions to the*

History of Starch and its Transformations. The experiments were conducted with well-washed and purified potato-starch. The transforming agent used was a cold, filtered, aqueous infusion of malt, made by acting upon 100 grms. of finely-divided pale malt with 250 c.c. of water. The total solid matter is deduced from the specific gravity, a constant divisor 3.86 being used. The starch granule consists of granulose and starch cellulose, the latter preponderating in the outer layers. Starch cellulose can be obtained by acting upon starch paste in the cold with unheated normal malt extract. The granulose rapidly dissolves, leaving most of the starch cellulose behind. When once separated in the insoluble form this substance cannot again be brought into aqueous solution without decomposition; it is also unattacked by malt extract. On boiling with water it is chiefly converted into soluble starch, which has probably no action on polarised light. The viscosity of starch paste varies much with the previous treatment of the starch. The specific rotatory power of starch paste equals 201.1° to 202.1° . The specific rotatory powers of granulose and soluble starch are identical. With potash, the specific rotatory power of granulose is 169.4° to 174.2° , and of soluble starch 182.6° . The maltose used was prepared by acting on starch with malt extract, and crystallisation from alcohol. Its specific rotatory power was 150.4° ; its cupric oxide reducing power was 61.0. By the action of dilute sulphuric acid a dextrose identical with that of invert sugar was obtained. The end product of the reaction had a specific rotatory power of 58.6° , and a cupric oxide reducing power of 100.2. Maltose is unacted upon by continued digestion with malt extract at 50° to 65° C. As the outer layer of the starch granule consists of insoluble starch cellulose, the granulose cannot be dissolved by any converting agent which is not diffusible, and which does not at the same time transform it into diffusible products; so the intact starch granule can be digested with malt extract, the transforming agent of which is colloidal, at ordinary temperature without action. If, however, the granules are ruptured by trituration with sharp quartz sand a rapid action is set up. Starch cellulose at first enters into solution, afterwards maltose and dextrin are formed. The action of malt extract on starch-paste is very rapid; in about three hours the specific rotatory power attains a maximum of 161.6° ; the cupric oxide reducing power also reaches a maximum at the same time, =40.7. The modifications of this reaction at elevated temperatures depend much more on the previous treatment of the malt extract than upon the actual degree of temperature at which the transformation is made; this fact has been noticed by O'Sullivan. The authors state in general terms that "if two different transformations of equal quantities of starch are brought about at different temperatures by equal quantities of a malt extract which has been heated for a few minutes to a point identical with or higher than the more elevated of the two temperatures of transformation, then the conversions will be similar in every respect." To investigate the various stages of the action more thoroughly it was necessary to be able to arrest the reaction at any desired moment. This was done by pouring the sample into a small flask containing a few centigrams of salicylic acid. The authors then give tables and curves illustrating the action of malt extract upon starch paste at various temperatures and under varying conditions. Throughout these transformations the specific rotatory power and the cupric oxide reducing power of the products agree closely with the supposition that maltose and dextrin are the only substances formed in the reaction. When the products of a transformation of starch having a united optical activity higher than 162.5° are treated with a little unheated malt extract at 50° to 60° the specific rotatory power falls at once to 162.5° ; this change takes place *per saltum* and not gradually. The most probable formula of soluble starch is $10(C_{12}H_{20}O_{10})$. The first action of the transforming agent of malt extract is the hydration of one (of the ten) $C_{12}H_{20}O_{10}$, and the consequent production of maltose, the nine left forming erythro-

dextrin: the remaining changes are given in the following table:—

	First power. Degrees.	Copper oxide reducing power.	Resulting dextrin.
Soluble starch	216.0	0	—
1	209.0	6.4	Erythro-dextrin, α
2	202.2	12.7	" β
3	195.4	18.9	Achroo-dextrin, α
4	188.7	25.2	" β
5	182.1	31.3	" γ
6	175.6	37.3	" δ
7	169.0	43.3	" ϵ
8	162.6	49.3	" ζ
9	156.3	55.1	" η
Maltose	150.0	61.0	—

The authors have established the existence of Nos. 2, 3, 4, and 8, and have indications of 5 and 6; No. 8 is the most stable. Dextrose is not a product of the action of malt extract upon starch, the hydration ceasing with the complete conversion into maltose. Diastase is a function of the coagulable albuminoids of malt extract, and does not consist of any particular principle. The albuminoids of barley resemble those of malt, but are less active. The albuminoids of barley precipitated above 60° are inactive as regards diastatic action. During germination the comparatively inactive albuminoids of barley have conferred upon them a large amount of potential energy. The growing yeast cell is capable, to a certain extent, of imitating the modifications induced by the living vegetable cell in the ordinary process of germination.

The next paper was read by R. WARINGTON, "*On the Determination of Nitric Acid by means of Indigo, with especial reference to Water Analysis.*" The indigo method possesses the advantages of great simplicity, speed, and delicacy. The results are, however, conditioned by many circumstances, which must be known before the method can be applied with accuracy. A solution of pure indigotin is much superior to the sulphindigotate of sodium usually employed, the former giving a pale colour when oxidised by nitric acid, and thus allowing a small excess of indigo to be readily seen. The estimation is made as follows:—20 c.c. of the water are mixed with the indigo solution, and pure oil of vitriol, equal in volume to the united water and indigo, suddenly added. The whole is then placed in a chloride of calcium bath at 140° till the reaction is completed. The quantity of indigo corresponding to the nitrate present is found by a series of approximating experiments. In weak solutions the quantity of indigo oxidised is not strictly proportional to the nitrate present. If a solution of nitre requires 10 c.c. of the indigo solution recommended in the paper a nitre solution of one-eighth of the strength will require only 1 c.c. The indigo must therefore be standardised with nitre solutions of graduated strengths, and a table of the value of the indigo scale constructed, by which subsequent analyses can be calculated. Some attention must be paid to the initial temperature of the solutions: a rise from 10° to 22° is attended with a diminution of 5 per cent in the indigo oxidised. Chlorides have some influence; a water containing 17.8 per million of nitrogen would yield 17.5 in the presence of much chloride, while a water containing 4.3 would give 4.5; the error is thus in opposite directions for weak and strong solutions. Nitrites cannot be determined by indigo; they oxidise less indigo than nitrates, and give no sharp reaction; they must be converted into nitrates by permanganate before employing indigo. Some kinds of organic matter greatly diminish the amount of indigo oxidised, and the soluble organic matter of soils affects the results in this manner. Analyses of waters by Frankland's method and the indigo process give closely concordant results.

The following papers were taken as read:—

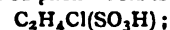
"*Notes on the Purple of the Ancients,*" by EDWARD SCHUNCK. This colour, which was extracted by th

ancients from various kinds of shell-fish, and applied to the dyeing of linen and woollen fabrics, has excited much interest from time to time. The author gives a *résumé* of previous work on the subject. Cole (*Phil. Trans.*, 1865) discovered a shell-fish in Somersetshire which yielded a purple dye. Reaumur in 1710 discovered a similar species (*Purpura lapillus*), Dutramel in 1736, and Bancroft in 1803, also worked at the subject. Their results may be summarised as follows:—1. The colour-producing secretion resembles pus, and is contained in a small whitish cyst under the shell close to the head of the animal. This pus-like matter when applied to white linen and exposed to sunlight changes from yellow, through light green, dark green, blue, to purplish-red or crimson, a strong odour resembling garlic or assafœtida being evolved. Daylight is essential; if kept in the dark the secretion remains unchanged for years, and then on exposure to light becomes coloured. Heat does not promote the change which proceeds *in vacuo* or in hydrogen or nitrogen gas. The colour resists the action of soap alkalies and most acids; it is destroyed by nitric acid and chlorine. A. and G. de Negri have recently obtained two colouring matters from a species of murex; one is blue and presents the characters of indigotin, the other is red, but its nature is not certain. The author has made many experiments with *Purpura lapillus*, which he obtained on rocks at low water near Hastings, live animals alone being used. The pale yellow secretion turns purple when exposed to light without being applied to linen. Boiling does not hinder the production of the colour. The author observed the change of colour under the microscope. The chromogen can be completely extracted by alcohol and ether from the pounded cysts, a golden solution being obtained, which becomes purple on exposure to light, a purple powder being ultimately precipitated, which is granular and crystalline. Hydrochloric acid produces a decomposition somewhat similar to that effected by sunlight, a purple colour being formed. The author worked up the cysts or veins from 400 animals, and obtained 7 milligrams of the purplish powder by exposure to sunlight. It was insoluble in water, alcohol, and ether; slightly soluble in boiling benzol and boiling glacial acetic acid; easily soluble in boiling aniline, the solution giving a broad absorption band between C and D. Heated between watch-glasses, a sublimate of crystals with metallic lustre showing at their edges a deep indigo-blue colour. The colouring matter dissolves in strong sulphuric acid to a purplish solution, showing a band between D and E. From these reactions the author concludes that the colouring matter belongs to an unknown member of the indigo-blue group, and proposes to call it Punicin. The liquid from which the action of sunlight had precipitated the purple powder contained no glucose.

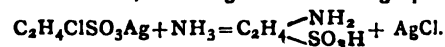
"On the Heat of Formation of Aniline, Picoline, Toluidine, Lutidine, Pyridine, Dipicoline, Pyrrol, Glycerine, and Furfural," by W. RAMSAY. In a recent paper on the volumes of liquids at their boiling-points the author remarked that some connection might exist between the ebullition volume of a liquid and the amount of heat evolved during its formation. The present paper contains an account of the method used to determine the heat evolved and the results obtained. The method employed was an indirect one; the amount of heat evolved by the combustion of the elements of a substance was calculated; the amount of heat evolved by the combustion of the substance was estimated; the difference between these numbers gives the heat absorbed in decomposing the compound; this difference is assumed to be equal to that evolved during its formation. Lewis Thomson's apparatus for determining the heating power of fuel was used. An oxidising mixture of 3 parts of potassium chlorate and one of potassium nitrate was used, with some sulphur added to promote combustion. The author discusses the probable errors of the estimation. The following table gives the principal results obtained:—

	Calories evolved.
Aniline	2747
Picoline	5753
Dipicoline	8084
Glycerine	-1364
Furfural	5985
Toluidine	1281
Lutidine	7184
Pyridine	7117
Pyrrol	4056

"On Ethylenic Chloro-sulphocyanide and its Oxidation into Ethylenic Chloro-sulphonic Acid," by J. W. JAMES. 66 grms. of potassium sulphocyanide were placed in a flask with about 250 c.c. of 98 per cent alcohol; after shaking, 100 grms. of chloro-bromide of ethylene prepared by Lössner's method were introduced, and the mixture heated on a water-bath; decomposition rapidly ensued. Potassium bromide and chloro-sulphocyanide of ethylene were formed; the latter substance after purification was obtained as a colourless oil, of a disagreeable smell resembling that of oil of mustard; it affects the eyes, does not excite sneezing, it boils at 202° to 203° C., its vapour burns with a violet coloured flame. Analysis gave the formula $C_2H_4Cl(CNS)$. By oxidation with fuming nitric acid, chloro-ethylen sulphonic acid is obtained—



the author has prepared and analysed the barium and silver salts. By heating the silver salt with ammonia in sealed tubes to 100° to 120° C. about 0.7 grm. of pure taurin was obtained, according to the following equation—



"On the Boiling-points of Certain Metals and Metallic Salts," by T. CARNELLY and W. CARLETON WILLIAMS. In a previous communication the authors described a simple method of measuring the temperature of ebullition of those bodies whose boiling-points are beyond the range of the mercurial thermometer, by observing whether certain salts whose melting-points are known melt on exposure to the vapour of the boiling substance. In the present paper they give results thus obtained:—Palmitic acid, 339° to 356°; stearic acid, 359° to 383°; selenium, 676° to 683°; tellurium dichloride, 327°; tetrachloride, 414°; tetrabromide, 414° to 427°; stannous chloride, 617° to 628°; bromide, 617° to 634°. By using pure metals instead of salts of known melting-points the following numbers were obtained:—Lead chloride, 861° to 1000°; cadmium chloride, 861° to 1000°. Glass tubes are rapidly corroded by molten potassium and sodium, iron tubes were therefore used. Sodium, 861° to 1000°; potassium, 719° to 731°; bismuth, 1090° to 1600°; lead, 1600° to 1800°; tin, 1600° to 1800°; antimony, 1090° to 1600°. The authors point out a curious relation between the melting and boiling point of Cl, Br, and I and those of S, Se, and Te; for the melting-points of the S, Se, and Te are respectively twice and the boiling-points three times as high as those of Cl, Br, and I, all being reckoned from the absolute zero -273° C. Thus:—

Melting points of—			
Chlorine	$198^\circ \times 2 = 396^\circ$	Sulphur melts	388°
Bromine	$251^\circ \times 2 = 502^\circ$	Selenium "	490
Iodine	$380^\circ \times 2 = 760^\circ$	Tellurium	755 to 773
Boiling points of—			
Chlorine	$240^\circ \times 3 = 720^\circ$	Sulphur boils	720°
Bromine	$318^\circ \times 3 = 954^\circ$	Selenium "	953
Iodine	$473^\circ \times 3 = 1419^\circ$	Tellurium "	?

The Spectrum of Didymium Nitrate.—Lawrence Smith and Lecoq de Boisbaudran.—Didymium nitrate, if neutral or slightly acid, gives an absorption spectrum almost identical with that of the chloride; the elementary rays of the bands are less distinct. An addition of nitric acid, however, produces certain important changes which the authors describe.—*Comptes Rendus*.

NOTICES OF BOOKS.

A Theoretical and Practical Treatise on the Manufacture of Sulphuric Acid and Alkali with the Collateral Branches. By GEORGE LUNGE, Ph.D., F.C.S. Vol. I. London: J. Van Voorst.

DR. LUNGE has been too long and too favourably known to the readers of the *CHEMICAL NEWS* to require any special introduction on our part as a preliminary to the notice of his *opus magnum*. The object of his treatise is three-fold. It aims in the first place at supplying a full and accurate account of all the raw materials, products, and residues of the chemical arts concerned. Secondly, it is addressed to students of technical chemistry, who will find here a description both practical and theoretical of the processes adopted in these manufactures. More especially, however, it seeks to give manufacturers, managers, &c., the best information upon the methods and the working plant which have come under the author's very extensive observation. Hence it is at once a record of practical experience and a careful and elaborate compilation, differing, however, from most works of the latter class by the circumstance that the various processes are not merely described but ably discussed, and that much unpublished information is here laid before the reader. English chemists and manufacturers will find here an account of certain processes and a statement of certain facts hitherto little known in this country, and will thus be enabled to compare point for point their own ways of working such as are current elsewhere.

Within the limits open to us anything like a full analysis of a work so rich in interesting matter is impossible, and we must confine ourselves to a notice of passages of especial interest. As a curious and important fact we find that, contrary to the usual rule for metals, lead is the more rapidly acted upon by sulphuric acid the purer it is. Small quantities of copper and antimony increase its resisting power, while traces of bismuth have an opposite action.

In the volumetric determination of free acid the author from his own experience recommends tropæolin (OO) as an indicator in place of litmus. He objects to oxalic acid for the preparation of normal acid for alkalimetric purposes; it is rarely pure, its crystalline water is uncertain, and in dilute solutions its stability is very imperfect. For the detection of nitric acid in sulphuric acid the author recommends carbazol, which, with traces of nitrogen oxides, yields an intense and permanent green colour.

Turning to the raw materials of the manufacture we still encounter the fact that now, just as was the case seven years ago, the sulphur deposits of Iceland and of Saba remain undeveloped.

The sources and the composition of the various kinds of pyrites now in the market are given at some length. As regards Norwegian ores the statement of H. A. Smith that arsenic is here present to the average amount of 1.649 is considered doubtful. The gross inaccuracy of the "Cornish assay" for copper ores is duly noticed. It is not creditable that such a process should still be recognised in a civilised country. In Chili it has long ago, we understand, been formally prohibited for commercial purposes.

Passing over the long and interesting account of the conversion of sulphur into gas, where almost every variety of kiln used successfully for different kinds of ore is described and figured, we come to the construction of the chambers. Here Dr. Lunge remarks that there is now little diversity of opinion on the question of form, cubical chambers being now very rarely laid out. The conclusions of H. A. Smith that sulphuric acid is formed at heights of 3 to 6, or at most 8 feet, the space above this being substantially wasted, is closely criticised. The statement that 94° C. is the normal temperature for sulphuric acid making is pronounced entirely unsupported

by experimental evidence. Smith's argument to show that the acid is almost solely formed at or near the bottom of the chamber is found to be fallacious. As a decisive experiment he had placed two collecting-dishes, one at 4 feet and the other at 16 feet from the chamber floor. After nine days he found, indeed, that the lower dish had collected much more acid than the upper, but he overlooked the fact that an uncovered dish placed near the bottom of the chamber receives not merely the acid formed immediately above it, but also all that is formed in the entire space directly over it, whence the experiment utterly fails to show where the acid is chiefly produced. Haas cleverly modified Smith's experiment by fixing dishes at different heights, placing a leaden cover at the height of a foot above each. The quantity of acid formed all over the chamber was then found almost equal. According to Bode a set of chambers of Smith's dimensions was actually erected at the Oker Works, but was found to yield acid not in the ratio of its area, as had been assumed, but of its cubical contents. Hasenclever, in opposing Smith, flies to the opposite extreme and recommends chambers 33 feet in height. Many years experience at the Oker Works have shown that "16½ feet width and height have been found the most suitable size." As regards the size of a set of chambers the author considers that the golden mean lies between 200,000 and 300,000 cubic feet. In smaller sets the same labour and auxiliary plant are needed, and the working is consequently relatively less economical. The total space required may range, it is considered, from 16 to 18 and even 20 cubic feet per lb. of sulphur charged. The Glover tower effects a saving in space, and in winter for the same space more pyrites can be burnt than in summer.

The best method of using the nitre, whether in the solid state or as free acid, is next taken into consideration. Here, speaking from memory, we should say that Dr. Lunge has to some extent modified the views which he formerly expressed. He now admits that "the best English works, all of which employ solid nitre, work with as small a consumption of it and as good a yield of vitriol as the best of the Continental works employing liquid nitric acid; and on the Continent as well many manufacturers work quite as well with solid nitre as their neighbours with liquid acid." He states that he would "in no case think of employing nitric acid except where it is made at the works for sale." Solution of nitrate, he considers, would combine every advantage, but it has not yet been successfully introduced.

From the points described as essential to the successful working of a set of chambers, we quote merely the recommendation so to regulate the draught that the burner-gas, if from brimstone may contain 11 per cent, and if from pyrites 8.5 per cent SO₂, and that the gas issuing at the end should retain 5, or better, 6 per cent of oxidation. In Germany, we may remark, the draught is very generally regulated according to the indications of gas-analysis, whilst in England it is still too often left to "rule of thumb."

Space does not permit us to enter into the discussion on the working of the Glover tower, one of the most interesting portions of the whole work. The author considers it proved both by general experience in England and by the direct numerical results obtained by Bode that "the Glover tower remains by far the cheapest apparatus for denitrating and concentrating; it saves all cooling contrivances, and is able to bring up the whole of the acid to 152°, nay, even sometimes up to 160° T. That against these advantages there is no set-off of a larger consumption of nitre has been mentioned previously."

We predict for this work a great success. To the best of our knowledge it is the most complete, practical, and thorough-going treatise on the subject to be met with in any language. Chemical manufacturers, managers, works' chemists, technical students, &c., will find it indispensable.

Improvements in Sulphuric Acid Manufacture. (Read at the General Meeting of the Institution of Civil Engineers of Ireland, April 2, 1879.) By W. G. STRYPE, C.E. Dublin: Forster and Co.

THE author begins with a complaint concerning the shortcomings of English chemico-technical literature, and criticises somewhat sharply the works of W. H. Smith, C. T. Kingzett, and A. G. and C. G. Lock, complaining that the last-mentioned "does not contain a single chemical or mathematical formula!" He next gives a sketch of an ideal treatise on the manufacture of sulphuric acid. He then enters upon a description, and takes the *CHEMICAL NEWS* to task for having, in a review of the Messrs. Lock's work, given an unfavourable account of Irish pyrites. He thinks it "hardly fair that statements should be made in the absence of proper enquiry." The reviewer, however, in characterising these pyrites as, "on an average, low in sulphur and very hard to burn reasonably clean" was guided by prolonged personal observation, made about twenty years ago in the Widnes district, and by very numerous analyses both of the green and burnt ore. If Mr. Strype will consult Dr. Lunge's recent work he will find there Irish ore described as "too hard and slaty," and not burning well, and as containing "only 30 to 35 per cent of sulphur."

On the important question of the supply of moisture to the chambers the author suggests that in preference to "pulverised water," moderately superheated steam should be used. Whether this suggestion has anywhere been put into successful practice we do not learn. The absorbing tower at the Wicklow Chemical Works is next described, but his account cannot be here introduced without the aid of the accompanying plates. The self-acting apparatus which he has devised for regulating the supply of air to the chambers seems to us a very valuable and desirable feature, and we hope to hear that its success has been confirmed by other observers.

CORRESPONDENCE.

INDUCTION DISTURBANCES IN TELEPHONE CIRCUITS.

To the Editor of the Chemical News.

SIR,—The report of my communication to the Physical Society on June 14, will, I fear, be as unintelligible to your readers as it is to myself. The method suggested is the employment of a shunt consisting of a cell containing platinum plates or wires in very dilute sulphuric acid. The induction currents on the line, having a high potential, escape through the shunt to earth, while some of the telephonic current passes to the telephone. In this way the induction currents are entirely removed, while the sound of the voice is only weakened.—I am, &c.,

HERBERT McLEOD.

Cooper's Hill, June 23, 1879.

[We inserted the report just as it was received from the official reporter of the Physical Society.—Ed. C.N.]

ON THE FORMULA OF THE SESQUIOXIDES.

To the Editor of the Chemical News.

SIR,—In a paper which I read before the Chemical Society in May, "On the Volumes of Liquids at their Boiling-points, obtainable from Unit-volumes of their Gases," I gave a determination of the specific gravity of chromyl dichloride, CrO_2Cl_2 , from which the following inferences may, I think, legitimately be drawn. Chromyl dichloride at its boiling-point has the specific gravity 1.7538. The "ebullition volume" of chromium necessarily depends on those assigned to oxygen and chlorine. From the above

number it can be shown that 22,326 volumes of gaseous chromyl dichloride, calculated to 0° and 760°, condense to form 88.4 volumes of the boiling liquid. From determinations by Kopp and others oxygen has been proved to have two values, 7.8 and 12.2, and chlorine the value 22.8. It is probable that in chromyl dichloride oxygen possesses both these values, consequently the value of chromium is easily calculated by the following equation:—

$$\begin{array}{ccccccc} 88.4 & - & (7.8 & + & 12.2 & + & 45.6) & = & 22.8 \\ \text{CrO}_2\text{Cl}_2 & & \text{O}_1 & & \text{O}_{11} & & \text{Cl}_2 & & \text{Cr.} \end{array}$$

The number assigned to chromium in determining the volume of boiling liquids containing that element, obtainable from 22,326 volumes of their gases, is therefore, 22.8. From this number the true formula of crystallised chromium oxide may perhaps be inferred. In Gmelin's "Handbook of Chemistry" its specific gravity is stated to be 5.21. Its molecular weight, 152, divided by that number gives its ebullition volume, 29. If the lowest value for oxygen be taken, viz., 7.8, the calculated volume of Cr_2O_3 is 74.3. Supposing that chromium oxide could be fused and boiled without decomposition, it is hardly conceivable that it should expand from 22 to 74, that is, increase its volume two and a half times. It appears, therefore, more probable that chromium oxide has the formula Cr_4O_6 , in which case it should have the volume 56; and it is not inconceivable that expansion from 56 to 74 would take place from the known infusibility of chromium oxide.

I had already written the above, but should not have published it had it not been for the very remarkable result since obtained by Victor and Carl Meyer, and published in the *Ber. der Deut. Chem. Gesell.*, xii., p. 1112, in determining the vapour-density of arsenic trioxide. At a temperature of 1560° C. the specific gravity of gaseous arsenious oxide was found equal to 13.80; the calculated vapour-density of As_4O_6 is 13.68. The coincidence between the two different methods of experiment and different trains of reasoning surely points to the necessity of a change in the formulæ of the sesquioxides.—I am, &c.,

WILLIAM RAMSAY.

Glasgow University, June 23, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 23. June 9, 1879.

Bases Derived from Aldol-ammonia.—A. Wurtz.—A description of tricrotonylen-amin and its nitrate, hydrochlorate, chloro-platinates, and chloro-aurates.

Increase of Albuminoid Matters in the Saliva of Albuminuria Patients.—M. Vulpian.—In one case 0.45 of mucin and 0.145 of albumin have been found in 1000 grms. of filtered saliva.

The Spectrum of Erbium Nitrate.—Lecoq de Boisbaudran.—Erbium nitrate, if neutral and moderately concentrated, gives an absorption spectrum not differing sensibly from that of the chloride. The presence of an excess of nitric acid produces very distinct changes, which are here particularised.

The Laws of Dispersion.—M. Mouton.—A mathematical paper, not capable of useful abstraction.

The Law of Stokes.—S. Lamansky.—The author concludes from his researches that the law of change in the refrangibility of light holds good perfectly in the general form in which it has been propounded by its author.

Absorption Spectra of Alizarin and Certain of its Tinctorial Derivatives.—A. Rosenstiehl.—On comparing the spectra of nitralizarin, of purpurin, and pseudo-purpurin with that of alizarin it is easy to follow the effects of the substitution of compound radicals for hydrogen. The violet extremity of the spectrum disappears and the lustre of the violet diminishes in proportion as the thickness of the coloured stratum augments. The red part not merely retains its brightness but grows broader. An effect still more marked is found on turning to the bisubstituted derivatives of alizarin. The spectrum of pseudo-purpurin, which contains in place of H the group CO_2H , is an instance. We see merely a large dark band, which extends from side to side of the ray E and which obliterates all the green portion. The effects of substitution in the molecule of alizarin tend towards the effacement of the details of the spectrum. In proportion as the solution of a colouring matter is seen in greater thickness it appears redder.

Dissociation of Ammonium Sulphide.—R. Engel and M. Moitessier.—The authors, in opposition to M. Sainte-Claire Deville, maintain that 2 vols. ammoniacal gas and 1 vol. hydrogen sulphide combine at common temperatures to form ammonium sulphide, 1 vol. ammonia remaining free. At elevated temperatures the product is rapidly dissociated, the gaseous mixture at 45° occupying 3 vols. and not 2, as stated by MM. Deville and Troost.

Action of Watery Vapour upon Carbonic Oxide in Presence of Platinum Wire Heated to Redness.—J. Coquillion.—The result is a mixture of carbonic acid and watery vapour.

Certain Derivatives of Methyl-eugenol.—M. Wassermann.—The compounds described are the dibromide of mono-bromated methyl-eugenol, methyl-eugetinic acid, mercurio-dimethyl-eugenol, and a product of the oxidation of methyl-eugetinic acid.

An Isomer of Angelic Acid, the Dimethyl-acrylic Acid.—E. Du villier.—The author admits the justice of M. Miller's reclamation, but points out that he has effected the synthesis of this acid by a novel process.

Localisation of Arsenic in the Brain.—O. Caillol de Poncy and C. Livon.—Arsenic seems to be substituted for the phosphorus of phospho-glyceric acid, thus producing an arsenio-glyceric acid. Lecithin then is modified so as to contain arsenic in place of phosphorus.

Hannoversche Monatsschrift wider die Nahrungsfalscher.
Vol. ii., No. 2, February, 1879.

Washing-Mixtures.—The author gives analyses of a number of washing-crystals, powders, &c. In one case common salt occurs to the extent of 24.6 per cent, in addition to 9.2 per cent sulphate of soda. In others sulphate of soda is found amounting respectively to 22.37 and 21.9 per cent. The author ventures the inference that the consumption of these articles in any town must vary inversely as the culture of the inhabitants.

American Association for the Advancement of Science.—Dr. H. Carrington Bolton informs us that the twenty-eighth meeting of this Association will be held at Saratoga, N.Y., commencing on Wednesday, August 27, 1879. The following is a list of the Officers of the Saratoga Meeting:—President, George F. Barker, of Philadelphia; Vice-President, Section A, S. P. Langley, of Alleghany; Vice-President, Section B, J. W. Powell, of Washington; Chairman of Permanent Subsection of Chemistry, Ira Remsen, Baltimore; Chairman of Permanent Subsection of Microscopy, Edward W. Morley, of Hudson; Permanent Secretary, F. W. Putnam, of Cambridge; General Secretary, George Little, of Atlanta; Secretary of Section A, John K. Rees, of St. Louis; Secretary of Section B, A. G. Wetherby, of Cincinnati; Treasurer, William S. Vaux, of Philadelphia.

MEETINGS FOR THE WEEK.

SATURDAY, 28th.—Physical, 3. "On a New Polariscopes," Prof. W. G. Adams. "On the Spirality of Magnetic Energy," Dr. R. C. Shettle. "On the Conjugate Positions of two Circular Coils," W. Grant. "On the Distribution of Heat in the Spectrum," Sir J. Conroy.

FRIDAY, July 4.—Geologists' Association, 8.

ERRATA.—P. 276, col. 2, line 27 from top, for $q(t+x)=q'(t'+y)$ read $q(t+x)=q'(t'+y)$. Line 37, for columns read column.

INSTITUTE OF CHEMISTRY.

The President has offered Two Prizes of £50 each for the two best original investigations involving Gas Analysis. These Prizes will be open to Associates, and to all persons (except Fellows of the Institute) who shall before the 31st December next have qualified for the Associateship in all respects short of passing the prescribed practical examination, and successful competition for these prizes will be accepted in lieu of such practical examination.—Further information may be obtained on application to the Secretary, Mr. C. E. GROVES, Somerset House Terrace, W.C.

THE YORKSHIRE COLLEGE.

The Worshipful Company of Clothworkers of the City of London having increased their Endowment of the Textile Industry Department of the College so as to include instruction in Dyeing, the Council of the College is prepared to receive applications for the post of Instructor; Stipend, £300 a year and half the Fees. Preference will be given to Candidates who are familiar with the processes of French and German Dyeing and with the Methods of Instruction pursued in the Continental Schools. Applications, accompanied by a statement of the Candidate's experience, together with copies of Testimonials, to be sent not later than August 11th to the Secretary of the Yorkshire College, from whom further information may be obtained. The work of instruction will begin in January, 1880.

Leeds, June 18th, 1879.

W. F. HUSBAND, Secretary.

TO PROFESSORS & EXPERIMENTALISTS.

W. STONE, Mathematical, Surveying, and Optical Instrument Maker, 44, Gloucester Street, Queen's Square, Bloomsbury, works out all kinds of difficult experimental and scientific work.

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Transparent Colours are measured by moving a single reflecting surface of white glass; the liquid comes in contact with glass only; and there is no error from dark meniscus. Precipitates are made and supported in a suspensory fluid; a black button is caused to descend through this and vanish, then measuring the relative weight of the precipitate.

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Analytical Chemists will find this a concise and yet complete book of reference for the isolation and examination of the active principles of drugs. Special appendix on the microscopic characters of the starches in food and drugs. Copious index and qualitative courses or resins, &c.

Published by W. BAXTER at the Office of the South London School of Pharmacy, Kennington Cross, S.E., and sold by Messrs. Simpkin and Marshall and Messrs. Baillière, Tindal, and Cox.

PATENTS.—Mr. Vaughan, F.C.S., British, Foreign, and Colonial PATENT AGENT. Special attention given to Inventions relating to Chemistry, Mining, and Metallurgy. "Guide to Inventors" Free by Post.—Offices, 67, Chancery Lane London, W.C., and 8, Houndgate Darlington.

Silicates of Soda and Potash in the state of Soluble glass, or in CONCENTRATED SOLUTION of first quality, suited for the manufacture of Soap and other purposes, supplied on best terms by W. GOSSAGE and Sons, Soap Works, Widnes, Lancashire.
London Agents, COSTE and Co., 19 and 20, Water Lane, Tower Street, E.C., who hold stock ready for delivery.

A Chemist (21), who has had practical experience in works, desires Re-engagement. Good references; no objection to go abroad.—For particulars, &c., address, X. T., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

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Analytical Chemist desires an Engagement. Good testimonials. London preferred.—Address, Analyst, 351, Middleton Road, Oldham.

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A French Gentleman, speaking English fluently, thoroughly conversant with English and French pharmacy, and an Associate of the Pharmaceutical Society, wishes a Re-engagement; London preferred. First class references, &c.—Address, Z.. Laboratory, 143, New Bond Street, W.

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Superior Iron Filter-Press for Sale, made to order and of extra quality, by Messrs. Needham and Kite. It contains ten chambers, each 19 x 21½ inches; is provided with a 3-inch gun-metal pump to work by hand or steam, and with fittings for washing and steaming, which can be used or not at discretion. The whole is quite equal to new, and is in perfect working order.—Address L. B. S., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

PHYSICAL SOCIETY.—The REPRINT of SIR CHARLES WHEATSTONE'S SCIENTIFIC PAPERS is now being issued to the Members of the Society. Any Member who, having paid his Subscription for the current Year, has not yet received a Copy, is requested to communicate with the Treasurer, Dr. ATKINSON, Portesbury Hill, Camberly, Surrey.

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London 18th June 1879

NOTICE—ARTIFICIAL ALIZARINE

WE the undersigned the Owners respectively of the Patents for the Manufacture of Alizarine

A D 1869 No. 1936
A D 1869 No. 1948
A D 1869 No. 3318

HEREBY GIVE NOTICE That we recently commenced Legal Proceedings against The Alizarine and Anthracene Company Limited for damages for the infringement of the above-mentioned Patents and for an Injunction to restrain the said Alizarine and Anthracene Company Limited from manufacturing selling or from offering for sale or disposing of or parting with any Artificial Alizarine or other Coloring Matters manufactured or prepared in contravention of our rights under the said Patents or either of them respectively And that after obtaining an interim injunction on the terms of our application we at the request of the said Alizarine and Anthracene Company Limited agreed to stay our Action upon the following terms

a. That the said Alizarine and Anthracene Company Limited should submit to a perpetual Injunction

b. That they should admit the validity of the said Patents

c. That they should pay damages for past infringements

d. That they should pay our Costs of the Legal Proceedings

A perpetual injunction against the Alizarine and Anthracene Company Limited have been granted and all proceedings in the Action have been stayed on the above terms

AND WE HEREBY GIVE FURTHER NOTICE that we recently were about to institute legal proceedings against Messrs. Arthur and Henshaw Drysalter and Oil Merchants in Glasgow and Matthew Clark and William Andrew Jamieson Drysalter and Oil Merchants Glasgow the individual partners of the said firm of Arthur and Henshaw to obtain interdict against their infringing the said Letters Patent or any of them and damages for infringements thereof by them in time past and that we have agreed to discontinue and discharge the said threatened proceedings upon certain conditions embodied in an Agreement entered into between us and them Under the said Agreement the said Messrs. Arthur and Henshaw and the said Matthew Clark and William Andrew Jamieson the individual partners of the said firm admit and acknowledge

a. That the said Letters Patent are valid in every respect

b. That they have infringed the said Letters Patent by importing vending and selling Artificial Alizarine manufactured according to one or other of the processes described in the Specifications respectively filed in connection with the said Letters Patent

c. That the Letters Patent of the 28th May 1869 No. 1642 granted to Franz Julius Bröner (therein named Julius Bröner) and Hermann Gutzkow are and always have been null and void

d. They undertake during the subsistence of our said Letters Patent not to import into or sell within the United Kingdom any Artificial Alizarine produced by any of the processes described in the Specifications of any of our said Letters Patent except with our consent in writing

WE LASTLY GIVE NOTICE that any person or persons infringing our above-mentioned Patents whether by *purshasing* (except from us or our Licensees) or *selling* importing or being concerned in importing or in anyway using Artificial Alizarine in the United Kingdom other than Alizarine made here by us or imported by us or our Licensees will be immediately proceeded against

A REWARD will be given to any person who will give the undersigned information sufficient to undertake legal proceedings against any such Infringers

BURT BOULTON and HAYWOOD

64 Cannon Street City London

Badische Anilin and Soda Fabrik

F. ENGELHORN AUGUST CLEMM

BISULPHIDE OF CARBON, CHLORIDE OF SULPHUR.

GAS PURIFICATION & CHEMICAL CO., LIMITED,
161, 162 163, PALMERSTON BUILDINGS, LONDON, E.C.

S. A. SADLER,

CLEVELAND CHEMICAL WORKS,

MIDDLESBROUGH;

Newfall Tar Works, Carlton;

Ammonia Works, Stockton-on-Tees;

and Stamshaw Chemical Works, Portmouth.

And also of the Furness Tar Products Co., Ulverston.

Manufacturer of Benzole, Toluole, Xylol,

Solvent and Burning Naphthas, Carbolic Acid and Disinfecting Powder, Refined Anthracene Naphthalene, Black Varnish, Refined Tar, Crude Liquid Ammonia, Coal-Tar, Pitch, Creosote, Grease, Sulphate of Ammonia, Pyroigneous Acid, Acetate of Lime, Wood Naphtha, Charcoal, &c., &c.

S. A. S. is always a buyer of Coal-Tar Naphthas, Crude Anthracene and all Tar Products.

All communications to be addressed to the offices at Middlesbrough.

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THE CHEMICAL NEWS.

VOLUME XL.

EDITED BY WILLIAM CROOKES, F.R.S., &c.

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NOTES ON SOME ANALYSES OF WATERS.

By Dr. T. L. PHIPSON, F.C.S. Lond.,
Member of the Chemical Society of Paris, and of the Royal Society
of Med. and Nat. Sciences of Brussels.

THE following notes of some analyses made by me during the last few years may prove interesting at a time like the present, when this particular subject is attracting so much attention. As I have stated in my letter in the CHEMICAL NEWS (vol. xxxix., p. 63), the first question to be solved by the analytical chemist is whether the water submitted to him is fit to drink or not; and a very long experience in this class of investigations is requisite to enable him to pronounce with any degree of certainty upon this important point. In the next place it is required to know whether the water will attack and dissolve lead; whether it will corrode iron pipes or boiler-plates, and so prove injurious to those who use it. Finally, a whole host of considerations crop up as to its fitness for household and manufacturing purposes, into which there is no need to enter here.

Much stress has of late years been laid upon the *organic matter* contained in waters, and it is well to bear in mind that although a large amount, such as 6 to 8 grains per gallon, for instance, may be looked upon with suspicion, certain waters that contain as much as this may be drunk with impunity, whilst others that contain considerably less are known to be exceedingly injurious, if not fatal.

The *crenate of ammonia*, to which I alluded as being present in many spring waters (CHEM. NEWS, vol. xxxix., p. 62), and is taken up by alcohol from the dried residue, does not appear to be at all hurtful when present to the extent of as much as 4 or 5 grains per gallon; whilst putrid organic matter, numerous *bacteria* and *micrococci*, and minute *white* fungoid growths, are sources of imminent danger.

But without further general remarks I shall content myself on this occasion by giving the following results, which, incomplete as they are, appear to me to possess a certain amount of interest, and may give some idea of the great variety of waters which, in the course of a comparatively short space of time, come under the notice of the analytical chemist:—

I. Well near Sleaford (Lincolnshire).

Water not quite clear, with an *alkaline reaction* and decidedly *saline taste*, well aerated; contains some minute *green algae*; gases dissolved are air and carbonic acid only.

Total residue, 169 grains per gallon imp.

Chloride of sodium		76.0	grs.
Carbonate of soda		44.0	"
Sulphate of soda		35.0	"
Sulphate of potash		2.0	"
Chloride of magnesium		1.5	"
Organic matter		2.0	"
Phosphoric acid and bromine		faint traces	"
Silica		1.0	grs.
Ferric oxide		0.5	"
Carbonate of lime		7.0	"

169 grs.

II. St. Anne's Well, Buxton (Derbyshire).

Total residue, 20 grains per gallon.

Mineral matters		18	grs.
Organic matter..		2	"

The mineral matter consists *chiefly* of carbonate of lime and chloride of sodium, with a little sulphate of lime, traces of iron and silica; in the spectroscopic traces of caesium and strontium, no lithium or rubidium; the *residue* has a saline taste. Beautifully clear and tasteless; gives off carbonic acid on boiling, and a slight deposit of carbonate of lime. This water is said to have a constant temperature of 80° to 82° F. I found the specific gravity at 60° F. in my laboratory in London 1.003. The fact that this water cures the gout is owing probably to its great purity, to its being drunk warm and in large quantities, and to the diet, country air, and exercise prescribed to the patients.

III. Well on Wimbledon Common (Surrey).

Total residue, 32 grains per gallon.

Mineral matter		26	grs.
Organic matter and nitric acid		6	"

Mineral matter *principally* carbonate and sulphate of lime, with a very much smaller amount of alkaline salts. Well aerated. *No Phosphoric Acid*. A single drop of a very dilute solution of permanganate gives a rose tint to 200 c.c., which persisted for several hours. This is an example of a good well-water which has been used for drinking for many years.

IV. Well in the Lower Bagshot Sand near Esher (Surrey).

This well is 40 feet deep, and is situated about 40 feet from a small cemetery. The water is beautifully bright, clear, and odourless. It attacks lead easily, and dissolves

it. It shows decided indications of nitrates and much chlorides.

Total residue, 58·8 grains per gallon.

Nitric acid and organic matter	7·0 grs.
Chloride of sodium	14·0 "
Sulphate, carbonates, &c.	37·8 "

A very deceitful water. Certainly impregnated, and likely to get worse. Sp. gr.=1·0032. A spring much further from the little cemetery gave—

Total residue, 24 grains per gallon.

Nitric acid and organic matter	3 grs.
Mineral matter	21 "

This water also dissolved lead readily.

V. A Yellow Water; Locality Unknown (S. of England.)

This water was sent to me because it was supposed to be ferruginous and that it might have valuable medical properties. The locality was not mentioned. Seen in bulk, is golden yellow; quite transparent; has no sediment; when boiled, remains clear, but gives off a strong *marshy odour*.

Total residue, 2 $\frac{1}{2}$ grains per gallon,

Consisting almost entirely of *ulmate of lime* and *ulmate of ammonia*, with a little carbonate of lime and traces of chlorides, &c. Animal charcoal takes out the yellow colour. This water bleaches permanganate solution rapidly and in large quantities.

VI. Well at the Midland Bank, Birmingham.

Many well-waters in the town of Birmingham are said to attack iron, pumps, boiler-plates, &c.; this is supposed to be in virtue of the nitrates they contain, and "to be due to galvanic action."

Total residue, 81·62 grains per gallon.

Dry mineral matter after calcination.. 58·71 grs.

Contains a *very large* amount of *nitrates* and *ammonia*.

A bad water, quite unfit for household use, and said to be destructive to metal work. It would require special experiments to decide upon the nature of the action by which such a water corrodes metals, and these have not been made.

VII. Well in an Artificial Manure Manufactory, near Southampton.

Total residue, 3320 grains per gallon.

Free sulphuric acid	1500 grs.
Phosphates and sulphates of lime, alkaline salts, &c... ..	1820 "

This water had been used for some time, both for drinking—occasionally for making tea—and for replenishing a small boiler, until its taste became very acid, and it was then sent to me for analysis.

VIII. Well at the Albany Barracks, London.

Total residue, 80 grains per gallon.

Organic matter and nitric acid	8 grs.
Mineral matters.. ..	72 "

Supposed to have caused an outbreak of typhoid (no details).

IX. Well near Huntington.

Water used for drinking. It was sought to ascertain whether it would be apt to produce calculus.

Total residue, 82 grains per gallon.

Sulphate of lime	36·89 grs.
Carbonate of lime	15·37 "
Chloride of sodium	16·00 "
Organic matter and nitric acid	5·00 "
Silica, magnesia, oxide of iron, &c... ..	8·74 "

Could not be proved unwholesome; would not produce

calculus, but might prove prejudicial to the cure of that disease.

X. Water from a Scullery Pump in Bolton Street, Piccadilly, W.

Total residue, 1024 grains per gallon.

Half-a-pint evaporated gave a drachm of yellow liquid, closely resembling urine in all its characters. Said to have been "used for drinking, and to have produced sickness and diarrhoea." By addition of ammonia this water gave a precipitate in which was found *abundance of phosphoric acid*.

XI. Wells at Putney, S.W.

The *total residue* varies from 38 to 120 grains, and even more; some, which yield from 38 to 48 grains, with 7 or 8 grains of organic matter and nitric acid, have been used for drinking for many years without any apparent inconvenience. Others, that yield over 60 grains of *total residue*, containing more than 10 grains of organic matter and nitric acid, are proscribed by the medical authorities. A well in this locality gave me 63 grains of *total residue*, with 14 grains of organic matter and nitric acid, in the winter of 1878-9; and another well, a few yards further on, was analysed by the late Dr. Dundas Thomson (in 1854, as I afterwards discovered), with precisely the same results: the figures were almost identical. This would appear to show that for the last twenty-five years the well-waters in that particular spot have not varied in composition, *though there is evidence of contamination*.

XII. Observations.

The depth of a well has no influence upon the quantity of residue its water leaves upon evaporation. Thus, I found the total residue yielded by the water of a shallow well in the grounds of the Countess of Leven, at Roehampton, to be 38 grains. This well is 14 feet deep. Another, nearly the same depth and not very far distant, gave 119 grains.

The purer a water is the better it dissolves lead (as I have shown in my paper read at the meeting of the British Association at Norwich in 1868), and many a good spring water has thus been spoiled by storing in lead cisterns, being supplied by lead pipes, or pumped through a leaden pump. Slate cisterns are preferable, but they should not be soldered at the joints by red-lead, as is usually done.

Many years ago I showed that *boiler deposits* in all parts of the world (with the exception of a few special cases and marine boilers) are due to *carbonate of lime precipitated by the expulsion of carbonic acid from the boiling water*. They consist almost entirely (over 90 per cent) of carbonate of lime, similar to the mineral aragonite, in whatever country they are examined, as I showed for the first time in 1868 (*Scientific Review*, January, 1868).

I have tested a very great number of spring- and well-waters for *phosphoric acid*, and have usually found it *entirely absent*. Its presence is *always a bad indication*.

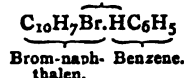
In the above analyses the microscope was invariably used for the examination of any residue deposited by the water on standing. In cases of sewage contamination through old-fashioned brick drains, it has more than once enabled me to detect dangerous waters by proving infiltration when the analysis was tolerably good. I am at present again occupied with some experiments on the organic matters of well-waters, and when these are completed I hope to give a brief account of them and of some tests to which a sample of water must be subjected in order to arrive at its true nature, and to be enabled to pronounce with certainty upon its fitness for drinking or other purposes.

Transmission of the Mineral Matters of the Soil into Vegetables.—M. Baudrimont has ascertained experimentally that recently precipitated tricalcic phosphate, calcic fluoride, ferric hydrate, hydrate silica, and aluminic hydrate are all soluble in an aqueous solution of carbonic acid.

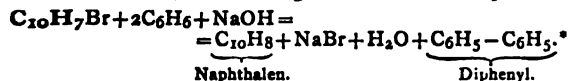
PRELIMINARY NOTE ON AN ATTEMPTED
SYNTHESIS OF THE HYDROCARBON PHENYL-
NAPHTHYL, OR PHENYL-NAPHTHALEN.

By WATSON SMITH, F.C.S., F.I.C.

In trying a series of experiments, last December, on the action at a red-heat of brom-naphthalen on naphthalen, both alone and in presence of soda-lime, by which dinaphthyl was pretty plentifully obtained (see *Journ. Chem. Soc.*, May, 1879, and *Ber. Deut. Chem. Ges.*, xii., No. 6, p. 674), the idea occurred to me that very possibly, by taking benzene instead of naphthalen, I might get the hydrocarbon phenyl-naphthyl or phenyl-naphthalen formed, according to the little scheme—



The proportions indicated by the desired reaction were taken, and the mixture was passed, first, alone through the red-hot tube; and secondly, over soda-lime, which had been arranged to form a layer in the tube. The monobrom-naphthalen was dissolved in the benzene, and the fluid was dropped into the tube by the arrangement described and sketched in *CHEMICAL NEWS* (vol. xxxix., p. 268). In the first case but little decomposition took place, even at the high temperature employed, and therefore but little hydrobromic acid was evolved. The result on subsequent examination was a small quantity of a resinous body boiling above 300° together with small quantities of other products (probably diphenyl, &c.). In the second case, a large quantity of naphthalen was formed, and came over. It is very probable, in presence of the soda-lime, the following reaction would take place:—



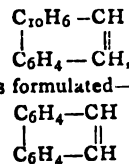
I now determined to try a reversal of the already mentioned mixture, taking one of brom-benzene and naphthalen; for it struck me this mixture would be preferable to the other, inasmuch as the boiling-points of the two constituents (brom-benzene and naphthalen) are not far apart, as in the former case, and I thought it not at all impossible that in like manner they might be more nearly on the same level with regard to what might be termed their dissociation-points, or at all events those points where the affinities of the H and Br are sufficiently weakened in their respective molecules ($\text{C}_{10}\text{H}_7\text{H}$ and $\text{C}_6\text{H}_5\text{Br}$), by the high temperature in the tube, that they shall incline to split off and effect a union, probably now lie much more nearly together than in the former case. Such a proximity is desirable as indicating, furthermore, a probable putting forth of powers of affinity in the nascent state, at about the same instant, and with something approaching the same degree of force. This was the consideration which led me to change the mixture as described, and I can now state that the results amply confirmed the stability of the theoretical ground taken in this case.

The mixture was passed through the red-hot tube drop by drop; hydrobromic acid was plentifully evolved, the amount of carbon separated was not great, and on subsequent examination of the black tarry-looking mass which had come over it was found to contain undecomposed naphthalen and brom-benzene, a very small quantity of benzene, diphenyl,—a body distilling over above 360°, but not at nearly so high a temperature as the dinaphthyls,—

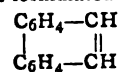
* I would argue from this that most probably the $\text{C}_{10}\text{H}_7\text{Br}$ (boiling at 277° is much more stable at the high temperature than the benzene (boiling at 80°), which is first decomposed. The nascent hydrogen then attacks the $\text{C}_{10}\text{H}_7\text{Br}$, diphenyl being at the same instant formed, and, in presence of NaOH , H_2O , NaBr , and C_{10}H_8 , are yielded—a reaction in detail taking place, and so obviating the formation of the desired phenyl-naphthalen.

and, finally, iso-dinaphthyl, with other yellow and red high-boiling products. It was found quite easy to separate the new body from the iso-dinaphthyl, since it is quite easily soluble in spirits of wine, in which iso-dinaphthyl is almost totally insoluble. The new substance crystallised from boiling dilute alcohol on cooling, in microscopic white crystals, agglomerated in tufts, and difficult to distinguish whether as needles or thin lanceolate plates. It was now placed on a filter, washed once or twice with cold dilute alcohol, and then water, and finally dried at 80°. The faint yellowish white mass was now submitted to sublimation, whereby beautiful little transparent plates were obtained, with a blue fluorescence resembling that of pure anthracen, and having a melting-point of 101° to 101½°. The odour of the vapours of this body resembles that of oranges. In the treatment with alcohol it was observed that one portion separated out from strong alcohol on cooling and standing over night, but the other portion refused to do so till after much more concentration. I think it very possible that these may turn out to be two isomeric bodies, and since two isomeric phenyl-naphthalens are possible it is quite probable they are here present. To this subject I am now giving the closest attention, and hope speedily to settle the question and be able to furnish final results.

The synthesis of phenyl-naphthalen, I need scarcely say,—since Graebe has so beautifully shown it to be the foundation, so to say, of chrysen, just as diphenyl is of phenanthren,—is a matter of no small interest in the territory of the aromatic compounds. Graebe finds (*Ber. Deut. Chem. Ges.*, xii., 1078), that, on heating chrysochinon with soda-lime, a hydrocarbon is formed having the composition of, and appearing to be no other than, phenyl-naphthalen, and so to chrysen he assigns the formula—



just as phenanthren is formulated—



on account of its similar relationship to diphenyl.

University Laboratory, Zurich,
June 28, 1879.

ON THE DIFFERENT ANILINE-BLACKS.*

By JUSTUS WOLFF.

(Concluded from vol. xxxix. p. 273.)

THE third series of aniline-blacks, of which the nigrosin which I invented is a link, we call the nigrosin series. This nigrosin was manufactured since 1863 in large quantities, and has been and is used for dyeing black and grey on wool, silk, and leather, especially in Germany, notwithstanding its high price (33/- per pound) at which it was sold during the first two years. A very large firm used it in great quantities for dyeing silk sunshades silver-grey, proving that it stands the action of light and air in no small degree. Its solution in alcohol in combination with varnish-producing oils and resins was used for black varnishing.

The first nigrosin was manufactured in 1863 by heating a mixture of 44 parts of aniline, 20 parts of stannous chloride, and 11 parts nitro-benzene in the first four hours to about 190°, and after that time to 220° to 230° for a sufficient length of time (taking about nine to twelve hours) until a sample drawn and poured into boiling water communicated to the latter a pure yellow colouration. At that point the unaltered aniline in the melt was driven off by a current of steam, the melt separated from the liquid

* All temperatures are given in Centigrades.

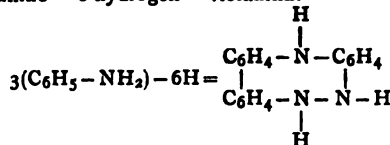
(produced by condensation of steam), dried, powdered, and sold as nigrosin. It is still in the market, and is used for the above-mentioned purposes.

I soon found out that the presence of stannous chloride was not essential for the production of that colouring matter, but that many other metallic salts would do the same, and even that it could be obtained without any metallic compounds at all, but only with aniline salt and nitro-benzene.

Inferring from that reaction that the nitro-benzene may be acting in this mixture as an oxidising agent, I used other oxidising compounds in its place—for example, arsenic acid—which, acting on aniline salt or a mixture of aniline and aniline salt in the proper proportions, produced also a fine nigrosin.

In trying to obtain this colouring matter in a water-soluble state I found that nigrosin made from impure aniline (containing toluydin) is not soluble in boiling water, but the latter dissolves a brown-yellow colouring matter; if the remainder is boiled two or three times more with water (slightly acidulated with hydrochloric acid), then it will dissolve some more nigrosin. The fourth extraction with slightly acidulated boiling water yields nigrosin in solution, and also all the following extractions. The fact that chlorine gas produces nigrosin with aniline in proper proportions and at a suitable temperature proves that it is in the first instance a product of dehydrogenation, and, following the process closely, we find that violanilin is formed in the first stage of the process, as the following formula represents:—

3 aniline — 6 hydrogen = violanilin.



The quantities of the dehydrogenators in the mixtures intended for the production of nigrosins being only sufficient to dehydrogenate and condense not quite the half of the weight of the aniline, we have, after this first stage of the process (the formation of violanilin), and when the dehydrogenation is finished, before us a mixture consisting principally of violanilin and aniline salt. By further heating this mixture to about 220° to 230° we see gradation-changes taking place in the colour of the mass or melt, from violet-blue to dark-blue, and later on to greenish black, whilst ammonia is formed.

Violanilin (prepared as pure as possible either from magenta-refuse or by the processes above described) heated with aniline salt (or mixtures of aniline and aniline salt) to 220° to 230° for a sufficient length of time yields water-soluble and insoluble nigrosin, besides ammonia.

Triphenyl-violanilin (the base of spirit-soluble indulin), or the spirit-soluble indulin itself, heated with aniline salt to 220° to 240° for a sufficient length of time, yields soluble and insoluble nigrosin, but no ammonia is formed.

Pure nigrosin is produced in the following way:—A mixture of 22 parts chlorhydride of chemically-pure aniline and 10 parts of chemically-pure syrupy arsenic acid (containing 70 per cent of dry acid) is heated for from four to five hours up to 190° in a glass or enamelled iron vessel, whilst well agitated from time to time. After that time the temperature is raised up to 220° to 230°, and kept at that heat for such a length of time that a drawn sample dissolves with a faint yellow colour in neutral boiling water. Then a strong solution containing a little more caustic soda than is sufficient to neutralise the acids in the melt is added, and the thus liberated unaltered aniline (besides some diphenylamin) is completely driven off by a current of steam. The remaining nigrosin base is separated from the alkaline solution, repeatedly washed with boiling water, and then treated with water acidulated with an excess of hydrochloric acid by boiling it for a sufficient length of time until nearly all is dissolved.

To the solutions obtained in that manner caustic is added, which separates the nigrosin base. The is collected on a filter, washed, and re-dissolved in slightly acidulated boiling water in order to obtain a strong solution of nigrosin. By addition of salt to that solution cooling nigrosin is precipitated, which, collected on a filter and well washed, is again dissolved in pure distilled water (the quantity of the water should be not quite sufficient to dissolve all the nigrosin whilst boiling), the solution filtered whilst boiling, separates nigrosin by cooling. This last operation, several times repeated, yields pure nigrosin.

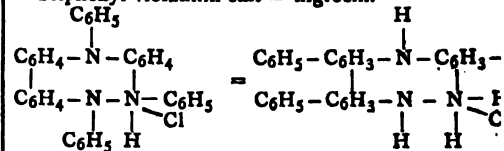
If the process has been conducted carefully, an aniline temperature kept in the limits given above, then nearly pure nigrosin, which, made from chemically-pure aniline, will be a blue colouring matter, whilst aniline containing toluydin in small quantities the shades of nigrosin are obtained in the manner given:—

100 parts of pure nigrosin dried for seventeen hours at 130° contain 7.2886 parts of hydrochloric acid. The average of four elementary analyses of the pure nigrosin base, dried at 130° to 140° for twelve hours, shows it consists of 86.368 per cent carbon, 5.387 per cent hydrogen, and 8.410 nitrogen. Consequently the formula for nigrosin base is $\text{C}_{36}\text{H}_{22}\text{N}_3$, and for the nigrosin $\text{C}_{36}\text{H}_{22}\text{N}_3\text{ClH}$. This is also the formula for triphenyl-violanilin. The conversion of triphenyl-violanilin in nigrosin base (both being isomeric) is therefore effected by an intramolecular change.

As the triphenyl-violanilin yields by dry distillation diphenylamin and aniline, and the nigrosin base does not yield any of them but substances belonging to the derivatives of di- and triphenylen-diamin it is easily inferred in the triphenyl-violanilin the hydrogens in the appendages are substituted by phenylen, C_6H_5 , whilst in the nigrosin the hydrogens of the nuclei are substituted by phenyl, $\text{C}_6\text{H}_5 - \text{C}_6\text{H}_5$.

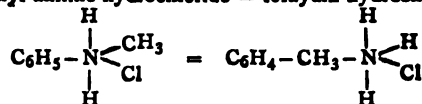
The triphenyl-violanilin salt is therefore converted into nigrosin by intramolecular transposition.

Triphenyl-violanilin salt = nigrosin.



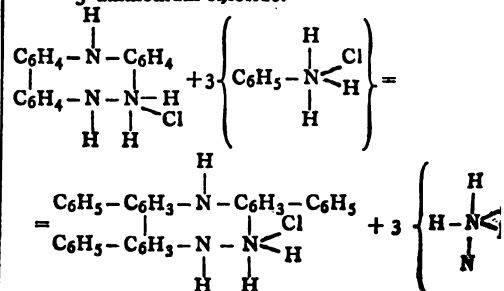
Hofmann published, in the *Ber. der Deut. Chem. Ges.* 1872, p. 720, a similar case in the conversion of aniline hydrochloride into toluydin hydrochloride by heating the former to 230° to 250° for a whole day.

Methyl-aniline hydrochloride = toluydin hydrochloride



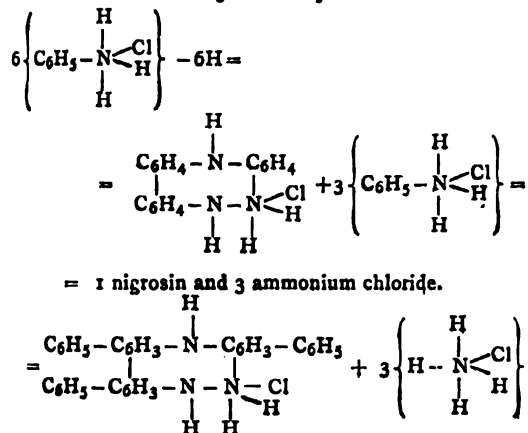
The formation of nigrosin from violanilin is represented in the following equation:—

1 violanilin salt + 3 aniline hydrochloride = nigrosin + 3 ammonium chloride.



And consequently the production of nigrosin from aniline hydrochloride—

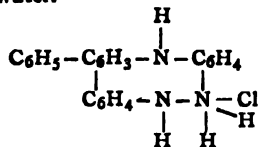
6 aniline hydrochloride — 6 hydrogen = 1 violanilin hydrochloride and 3 aniline hydrochloride.



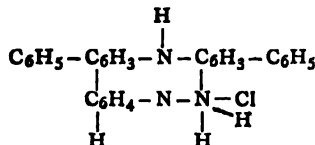
In the formation of indulin, as well as in that of nigrosin, a phenylation takes place. In the first case the nitrogen of the aniline acts as a triad, whilst in the violanilin the nitrogen of two of the appendants are triads, and that of the third appendant acts, being a compound with an organic acid at a low temperature as a pentad, which at higher temperatures starts dissociating and becoming triadic. Simultaneously it will be acted upon by the aniline triadic nitrogen, giving away its hydrogen and replacing it with the nucleus of the aniline; after that is done the other appendants are acted upon in the same way.

In the formation of nigrosin the nitrogen in the aniline and one, or also consecutively two, and three of the nitrogens in the violanilin act as pentads in combination with an inorganic acid which will not dissociate but at a very high temperature, and before this is reached the nuclei— C_6H_4 —of the violanilin on the one part, and C_6H_5 of the aniline on the other part, will act one on another, and nucleus substitutions will take place whilst ammonia is formed. Also in the conversion of triphenyl-violanilin into nigrosin the intra-molecular transposition creating this conversion is caused by the action of nitrogen as a pentad in the triphenyl-violanilin hydrochloride molecule.

The same may be said of triphenyl-mauvanilin, which, by the action of aniline hydrochloride, is converted into the brownish black nigrosin. Of course there are also nigrosins in which instead of three only two or one substitution has taken place, and such are insoluble in neutral and acidulated water.



And—



It is very probable that by the action of aniline hydrochloride on aniline-blue (triphenyl-rosanilin) at a temperature of 220° to 240°, by which De Laire's aniline-brown is formed, an analogous reaction takes place. This brown is also produced by the action of aniline hydrochloride on magenta and violet, analogous to the

formation of nigrosin from violanilin; and also analogous to the formation of nigrosin. Furthermore, it can be produced by the action of a dehydrogenator on an excess of a mixture of one molecule aniline hydrochloride and two molecules of toluydin hydrochloride. Here we meet with quite a new series of colouring matters, viz., those which can be obtained by simple intra-molecular transposition from other colouring matters caused by the action of its nitrogens in a pentadic state. These new colouring matters can also be obtained by dehydrogenation and subsequent phenylation of aniline, or mixtures of aniline and toluydin with pentadic nitrogens.

It is to be supposed that many reactions of this kind may occur, and they may be generalised as follows:—

I. By causing one or more of the constituent parts of a molecule to act on the other parts with another atomicity than they acted with in producing the original molecule, the latter can be converted into an isomeric compound (with different positions of its constituents), viz., intra-molecular transposition can take place.

II. By causing one or more of the constituent parts of a molecule to act at different times with different atomivities on the other parts, isomeric molecules with different positions of their constituents can be produced.

The pure blue nigrosin, whose production from chemically pure aniline salt has been given above, dissolves with a dull blue colour in neutral water, turning brighter and greener by addition of acid (hydrochloric acid principally). Its solutions in acidulated water possess a remarkable blood-red fluorescence, increasing by addition of more acid up to a certain proportion, and all the blue and black nigrosins possess this remarkable quality more or less.

This fluorescence is sometimes so strong that in a solution of blue or black nigrosin, which contains only such a small quantity that by looking through this solution no colouration is perceivable, it reflects a brownish red tint, especially before a black background, and the sun's rays falling upon it directly, it then looks as if very small particles of metallic copper were suspended in the liquid and moving about.

The slightly acidulated solutions in water dye wool, silk, China-grass, cotton, and certain other fibres (at about 80°) blue, the dye going on the fibre very slowly. If the bath is strong enough it will dye wool, silk, China-grass, and certain other fibres a fine blue-black (which, when deep enough, will stand light, air, and soap, but not the fulling process) in about one to one and a half hours without impairing the natural gloss of the different fibres. The same is the case with the other blue and black nigrosins; as all the nigrosins go on very slowly, they are especially adapted for dyeing yarns and goods evenly.

The following mixtures, treated in the same manner as described with that of aniline salt and arsenic acid, produce the different shades of the blue and black nigrosins:— 60 parts of hydrochloride of chemically pure aniline and 10 parts chemically pure nitro-benzene yield a dark blue dyeing nigrosin. The same mixture with 1 part cupric or cupreous chloride added to it yields a very fine deep blue-black. 60 parts of aniline hydrochloride (the aniline containing 2 per cent of toluydin) and 10 parts nitro-benzene (made of benzene containing 2 per cent of toluen) yields a blue-black dyeing colouring matter, which, by addition of certain metallic compounds (as, for example, cupric chloride, &c.), is considerably deepened.

By application of nitrobenzene in the nigrosin process other colouring matters besides nigrosin are formed in larger proportions than by application of certain other dehydrogenators, deriving from the action of intermediate products of decomposition of nitrobenzene, and remaining to the greatest part insoluble in neutral and acidulated water. By increasing the proportions of toluydin in the mixtures for nigrosin the black goes more and more over in jet-black and brownish black, which shades, by addition of copper and certain other metallic salts, are rendered deeper. Of course a considerable practice and a great

deal of experience is needed for the production of the different shades and qualities of the blue and black water-soluble nigrosins (patented in England, France, and the United States in 1876), as in this process the regulation of temperature at certain periods is of the greatest importance, for, if not carefully watched, a considerable quantity of by-products will be formed.

The nigrosins are slightly soluble in weak alkaline solutions by boiling. They dissolve easily in benzene, petroleum, and certain oils, especially when alkaline, with a bright purple colour, and when sour with a fine green-blue shade. Oxidising agents convert the blue and black nigrosins (especially when dyed in lighter shades on the fibre) into dull and reddish grey violets. Reducing agents render them colourless, forming leuco-nigrosins. Very remarkable is the resistance of the blue and black nigrosins against strong nitric acid, which, when dyed deep black, will, with nitric acid of even 1.5 sp. gr., only change to a very deep, nearly black-green after many hours of action, and then when the acid is washed off, the black does not seem to be changed at all. By this resistance against the action of strongest nitric acid, the blue and black nigrosins distinguish themselves from all other colouring matters.

Therefore, it is very probable that the chemical constitution of both series may be typically analogous, and thus the formulæ III., IV., and V. given by me to the Lightfoot black gain more probability, showing that both series have at least in common the nucleus substitution or connection of nuclei.

Substitution in Lightfoot Black.



Substitution in Nigrosin.



Further investigations may prove the analogy of both series—the one which I described by the formulæ III., IV., and V., and that of nigrosin.

The purer the aniline of which both are produced the bluer the shades of the blacks, and the blue and black shades of both are deepened by addition of certain metallic compounds. The Lightfoot-black turns green with acids, and the solutions of nigrosin also turn green with acids.

The application of the different nigrosins in dyeing is

the same for all, and like that given for the blue nigrosin. Nigrosin can also be used for printing blue, grey, and black on silk and wool in several ways, of which one is, to dissolve it in aniline or aniline salt, thickening the solution, and then to steam, wash, and dry. A solution of nigrosin in glycerin can also be used. Of course the insoluble and also soluble blue and black nigrosins can be converted with oil of vitriol into their respective mono-, di-, tri-, and tetra-sulphonic acids and their salts, which of course will improve the insoluble nigrosins and make them marketable, but not so the soluble nigrosins. The price of the nigrosins is not low enough that they could compete with the cheaper black colouring matters, but the beauty of their shades and their easy application, combined with resistance against light, air, and soap (except in the fulling process), will secure them a good market, especially for dyeing the more costly wool fabrics, China-grass, and above all silk. It is especially adapted for dyeing the last-named material a fine black without increasing its weight more than a few per cents, and being easily applied it could help to do away to some extent with the contemptible custom of weighting black silk fabrics. And, indeed, there should be put a stop to this pernicious practice, which reduces the famous durability of the silk fibre to its one-hundredth part by the corrosive action on the fibre of the materials used for weighting. The public pays the full price of silk for a material which contains only one-third of it (and that in a deteriorated state), and two-thirds of substances which have very little value compared with that of silk, and spoil it too. I know English manufacturers who send their silk to France for being dyed black and have it sent back for working it up, only for that reason, because in France they are able to increase the weight of silk whilst dyeing it black, much more than they can do in England; and therefore the silk dyed black in France will be cheaper by far than that in England, even after paying the freight to France and back. The result of such practice is a beautiful black silk fabric, changing into rags remarkably quick in the possession of the buyer.

Of course much has been said already authoritatively against this fraudulent adulteration, but it will not have any effect until numerous analyses of black silk fabrics have been made and their results published with the names of the respective firms manufacturing or selling them, and it would be a meritorious undertaking to organise in this manner a crusade against such offences.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

MAY, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

		Appearance in 2 foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	As Sulphuric hydride.	Hardness on Clark's Scale.	
			Saline.	Organic.								Before Boiling.	After Boiling.
<i>Thames Water Companies.</i>													
Grand Junction	Clear	0.000	0.006	0.141	0.067	21.11	7.630	0.790	1.224	1.53	14.3	3.7	
West Middlesex	Clear	0.000	0.006	0.135	0.070	21.00	8.170	0.860	1.152	1.66	14.3	3.7	
Southwark and Vauxhall	Clear	0.000	0.006	0.111	0.069	21.80	8.060	0.612	1.152	2.03	13.2	3.3	
Chelsea	Clear	0.000	0.006	0.135	0.058	20.20	7.280	0.540	1.224	2.00	14.3	3.3	
Lambeth	Clear	0.000	0.007	0.135	0.036	20.90	7.700	0.756	1.152	1.53	14.3	4.6	
<i>Other Companies.</i>													
Kent	Clear	0.000	0.004	0.295	0.004	20.70	10.410	0.936	1.512	3.03	17.6	6.0	
New River	Clear	0.000	0.004	0.135	0.030	20.70	7.590	0.468	1.152	1.16	13.7	3.3	
East London	Clear	0.000	0.006	0.105	0.052	23.00	7.840	0.840	1.296	2.07	14.7	4.6	

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

PROCEEDINGS OF SOCIETIES.

PHYSICAL SOCIETY.

June 26, 1879.

Earl Rosse in the Chair.

AN extra meeting of this Society was held on the above date at Cooper's Hill Indian Engineering College on the invitation of Col. Chesney, R.E.

Prof. UNWIN, of the College, read a paper on "*Experiments Relating to the Friction of Fluids on Solid Surfaces against which they rub.*" It has long been known that a board dragged through water suffers a resistance varying in some way as the square of the velocity, that a stream has a uniform motion at such a velocity that the component of the weight of the water down its inclined bed is balanced by the frictional drag on the bottom. The fluid in the neighbourhood of the stream is known not to move as a solid mass, the centre moving faster than the sides, and the different fluid layers rub against each other. The adhesion of the fluid to the solid, against which it moves, also gives rise to a sliding or shearing action. Our knowledge of the subject has hitherto been gained from observations on pipes, streams, and from the experiments of the late Mr. Froude with a plank of wood drawn through the water of a canal. It is desirable to have a set of laboratory experiments, however, as the conditions can be varied more than can be done by such methods, and for this purpose the author had designed a special apparatus. In Mr. Froude's experiments there was a practically unlimited mass of water and a definitely limited extent of solid surface; and his results are not free from certain anomalies. The author thought it might be instructive to try the other case of a limited mass of water, and a virtually unlimited surface; a disk in rotation gives such a surface. In some respects a cylinder would (as suggested by Prof. Ayrton) be the simplest to treat theoretically, but there are experimental difficulties in its way. The apparatus of the author consists of a metal disk rotated on a vertical axis in a vessel of water; and the problem is to determine its resistance to rotation, since this will be equivalent to the water friction upon it. Within the outer vessel is placed a thin copper chamber, the diameter of which is unalterable but the depth is variable at pleasure. The disk is placed concentrically inside this chamber, where there are two cheese-shaped masses of water, one above and one below the disk, which are dragged into rotation next the disk and retarded next the sides of the pan. The couple required to rotate the disk is equal to the couple exerted by the disk or the fluid when the motion is uniform. Hence the tendency of the chamber to rotate is measured by suspending the latter from three wires in a manner similar to the bifilar suspension of magnets. An index marks whether it rotates or not on a graduated scale, and a weight suspended by a cord measures the force required to keep the index at zero. Let M be the moment of the fractional resistance of the disk; N the number of revolutions per second. Then $M = CN^x$, where C and x are constants. The author has obtained a number of results, which are, however, not yet ready for publication. He mentioned, however, that a rough cast-iron disk has a frictional resistance almost exactly as the square of the velocity; whereas a turned brass disk gave a value of x decidedly less than 2. The resistance is a little greater when the mass of water is larger. These results were calculated for a speed of 10 feet per second. The author hopes to try the effect of temperature, &c., on fluid friction, and viscous as well as thin fluids.

Prof. UNWIN also exhibited a piece of apparatus with which he hopes to study the stress of rivetted plates under shear by means of elastic substances, such as caout-

chouc. He purposes to stretch the rivetted caoutchouc and photograph the appearance of stress-lines upon it.

Lieut. G. S. CLARK, R.E., explained the process invented by Prof. McLeod and himself for determining the absolute pitch of tuning-forks. Unlike other methods this is an optical one, and consists in arranging the tuning-fork to vibrate in front of a rotating drum whose periphery is marked with dots or fine lines at equal intervals. A microscope was arranged to comprise in its field of view the edge of the fork and the several of the intervals on the drum, so that when the drum was rotated at a rate which made the speed of an interval equal to the period of the fork, a set of prominences or waves, in width equal to an interval, were visible, the body of the wave being caused by the advance and recession of the fork in its vibration. The rotation of the drum was regulated by an air regulator devised by Prof. Unwin, the observer himself quickening or slowing the drum so as to keep the prominences steady. The time was got by an electric clock designed by Prof. McLeod. An aniline glass pen was used to mark the beginning and end of the periods of observation on the drum. A counter was also employed to give the number of revolutions. The pen and counter were actuated by electricity through the medium of a key. In these experiments a König fork, giving 256 vibrations per second, correct at 16.1°C. , was tested and found to give 256.2966 vibrations per second. Frequent bowing did not alter the phase. Fixing the fork rigidly, as in a vice, did so. The temperature coefficient for König's forks (0.00011 for each degree Centigrade) was confirmed by these experiments. Forks of different octaves were compared; audible beats could be counted; and modifications of Lesage's figures seen. This optical method is preferable to audible ones, because of its independence of the ear and the fact that nothing is attached to the fork itself.

Prof. GUTHRIE enquired if the periods of the forks had been found to alter through use or magnetisation.

THE AUTHOR said that he had not yet tested these points.

Prof. McLEOD instanced an old König fork which was correct at 16.1°C. , requiring now a temperature of 25°C. to make it so.

Lord Rosse suggested the use of the regulators employed with equatorial clocks to keep the rotation of the drum steady.

Capt. ABNEY enquired if a difference of vibration had been detected between the beginning and end of a series of observations. None had been certainly observed.

Prof. McLEOD then described an electric clock used in the foregoing experiments. A zinc and steel compensating pendulum moved by its own gravity; but at each beat made and broke a battery circuit by means of two bent springs, one on either side. The current passing through an electro-magnet detained a bent lever until the pendulum swung to the other contact. By this contrivance time was marked. Prof. McLeod found that the platinum contacts frequently stuck together in these experiments; but this defect had been cured by the use of a liquid shunt of dilute sulphuric acid, which destroyed the extra current. This remedy had been suggested to him by Lord Rayleigh. Prof. McLeod demonstrated the complete success of this device, which acts as well as a condenser shunt. He had also observed a curious effect with these liquid shunts, which as yet he could not explain. Two shunts, having the same acid in both, were employed, one shunting the extra current from four Daniell cells and one that from two Daniell cells. The first showed evolution of H and O gas, the platinum electrodes being unaffected. The second showed no evolution of gas, but one platinum plate was dissolved away and deposited in a black powder on the other. He also exhibited a new cell formed of zinc and mercury plates, with zinc iodide solution and mercurous chloride salt. Red iodide of mercury is formed at the negative electrode. The E.M.F. is seven-tenths of a Daniell cell, but the interval resistance

very low and the cell very constant, while there is no local action.

Prof. GUTHRIE suggested that as the extra current was really a succession of sparks the platinum might be carried bodily over from one electrode to the other.

Mr. F. H. VARLEY stated that Mr. F. Higgins had observed a similar effect with carbon electrodes in a voltameter, one carbon falling away into a fine powder, and due perhaps to the disintegrating action of liberated gases. He had also himself seen a platinum wire in contact with a carbon one eaten thin and drawn into very fine silky pens, while the carbon was stained blue, although the current passing was of low tension.

Mr. CHANDLER ROBERTS suggested that perhaps a hydride of platinum was formed in the case mentioned by Prof. McLeod.

Prof. GUTHRIE suggested experiments with fluorescent liquid shunts in the dark.

Mr. J. W. CLARK then described some experiments on the surface tension of sulphurous anhydride sealed in a capillary tube within a second tube containing the same substance. He found that at low temperatures the level of the liquid is lower in the narrow than in the wide tube. As the temperature rises the meniscus in the narrow tube descends, till at about 156° F. it is level with that of the wider tube, both surfaces being slightly concave. About this temperature the surfaces become plane then concave, the level in the wide tube becoming higher than that in the narrow one. These experiments are being continued, and Mr. Clark's other results will be communicated to the Society later on.

Prof. GUTHRIE proposed a vote of thanks to Colonel Chesney.

NOTICES OF BOOKS.

Note sur la Déphosphorisation au convertisseur Bessemer et sur la Déphosphoration des Fontes. Par M. POURCEL, Ingenieur Chef des Acieries de Terrenoire. St. Etienne: Théolier Frères. 1879.

THIS is a reprint of a Paper read by M. Pourcel at the meeting of the Société de l'Industrie Minérale, held on the 7th June last. After describing the rise and progress of the Thomas and Gilchrist method of dephosphorising iron, M. Pourcel gives an interesting account of his visit to Messrs. Bolckow and Vaughan's works, at Eston, in May last, where the process was carried out in his presence. The method of working in no way differed from that described by Messrs. Thomas and Gilchrist, in their Paper read before the Iron and Steel Institute in May last, and republished in the CHEMICAL NEWS (vol. xxxix., p. 219). The resulting metal contained—

C	..	..	..	..	..	0.171
Mn	..	..	..	..	..	0.160
P	..	..	..	..	..	0.223
S	..	..	..	..	..	0.037
Si	..	..	..	..	..	traces.

The crude iron used was Cleveland pig, containing 1.8 of P, 3 of Si, and 3.2 of C. The addition of 10 per cent of spiegeleisen at the end of the operation increased the quantity of P from 0.14 per cent to 0.223.

From his observations on this occasion M. Pourcel draws two conclusions. The first is, that the neutralisation of the silicic acid produced by the oxidation of the silicon in the crude pig is fully effected by the addition of a mixture of lime and iron oxide; and secondly, that overblowing, by prolonging the oxidising action, determines the scorification of the iron phosphide. The neutralisation of the silicic acid, adds M. Pourcel, gives rise to a large quantity of slag, which occupies a considerable space in the converter, diminishing its size from one-quarter to third. Hence, with very silicious pig, this change

operated upon falls from 8 or 9 to 6 or 7 tons. The "overblowing" increases the proportion of ferric peroxide dissolved in the bulk of metal, and consequently necessitates the addition of a considerable quantity of manganese—as much, indeed, as 2 or 3 per cent, if we want to avoid all danger of honeycombing. In the operations as carried out at Eston, M. Pourcel thinks that the "overblowing" was not kept up sufficiently long, because the metal of the last test, taken just before the addition of the spiegeleisen, contained 0.14 per cent of P, and that the manganese added in the spiegeleisen—1.7 per cent—was not sufficient, only traces of it being found in the resulting metal. The other defects in the process, as pointed out by M. Pourcel, are the several stoppages necessitated by having to take tests at different points of the operation, the uncertainty of knowing when the metal is sufficiently dephosphorised, and the explosions which result from the addition of the spiegeleisen—defects which the author looks on as being inherent at present, but which he hopes to see disappear in the course of working. Another very just remark which he makes is that the addition of spiegeleisen adds to the amount of P contained in the finished metal, not only from containing certain proportions of the very plague we are trying to get rid of, but by causing the reductions of part of the phosphate of iron in the slag owing to the liberation of large quantities of CO. The production of CO must therefore be reduced to a minimum, while the oxidising action of the air must be increased so that the P may not only pass into the slag, but remain fixed in it. M. Pourcel seems to think that if the P could be made to enter into the slag in the form of calcic phosphate, instead of iron phosphate, it would not be so liable to be reduced by the CO.

M. Pourcel recommends that the fumes which escape from the converter during the process of "overblowing" should be analysed; he thinks the greater portion of the iron phosphate is thus carried off. With respect to the part played by the lime in the process, M. Pourcel is at variance with most of the speakers at the meetings of the Iron and Steel Institute held in May last. The lime, he says, having played an active part of a neutralising base in the first part of the operation, thereby preserving the lining of the furnace from the corroding action of the silica, does not play a passive part during the operation of "overblowing." Calcium phosphate is a perfectly infusible body, the formation of which is not caused in this case by the action of the affinities at work, whether the lime is in the free condition or combined with the silica. M. Pourcel also combats the assertion of M. Gruner, who asserts that iron phosphide will not scorify until the carbon is reduced to the minimum. "How is it," asks M. Pourcel, "that Mr. I. Lowthian Bell, at the Clarence Works, and Dr. Bender, at Krupp's Works, succeed in dephosphorising their crude iron without diminishing the carbon contained in it? It is only necessary to quote the numerous analyses which prove that phosphorus disappears in direct proportion with the silicon." We learn, also, that Dr. Bender has adopted the Bell system at Essen, using a Pernot-Martin furnace, with iron sole and sides lined with rich manganiferous oxides. For several other interesting questions discussed by M. Pourcel we must refer the reader to his Paper.

The prime conclusion which M. Pourcel draws from his investigations is that the dephosphorisation of iron in the converter is an accomplished fact, and that the practical difficulties in the way of its industrial application may be surmounted by an attentive examination of the chemical phenomena which occur during the various operations.

Harmozein: a New Chemical Round Game, with Instructions for Playing it. By T. H. DAVIS, formerly Assistant at the Royal College of Chemistry. Manchester: J. Woolley, Sons, and Co.

HARMOZEIN, the name of which is presumably borrowed from the Greek *harmozo*, I sort, or arrange in order, is a

new scientific round game, which bears the same relation to chemical science that the much vaunted German *Kriegs Spiel* does to military strategy. Harnoezin must not be confounded with the numerous so-called instructive card games by which children are supposed to be deluded into the study of the 'ologies and 'ographies under the guise of innocent gambling. The instruction conveyed by these appliances generally consists in the acquirement of a mass of words, more or less hard; but any one who plays at Harnoezin must first of all have a fair knowledge of the theory of atomic quantivalence, which he will most certainly increase in proportion as he gains skill in the game.

The game consists of some 250 cardboard discs or counters, upon one side of which a chemical symbol is printed, with the quantivalence marked in the usual way, $\text{Cl}_{0.11}\text{Br}_{1.11}$, &c., the other side being left blank. The pieces are spread on the table blank side uppermost, and the game may be played by any number of players, each of whom selects from 10 to 20 pieces. The object of the game is to get rid of as many pieces as possible, each player being allowed to put down a formula as well as to add to those already on the board. Let us suppose, for instance, that the first player puts down COO, the second player, always provided he has the counters, puts down NaH and O, thus forming the complete formula of hydric sodic carbonate. He may then put down a formula of his own, let us say KCl, which no one can add to, as the monad K satisfies the monad Cl. He thus gets rid of four counters, whereas the first player only got rid of three.

Mr. Davis's game is sufficiently ingenious and interesting, but it might be improved. One of its objects being to teach the quantivalence of the elements, it seems strange that the pieces should be inscribed with the figures denoting it on the face. The quantivalence marks ought merely to be on the back, where the player is not supposed to see them. On the pieces denoting certain elements, such as iron, there is only one quantivalence mark, VI., but what is to be done when iron is a dyad, for instance. It would have been as well if the pieces denoting what Dr. Frankland calls polyad elements had been marked with their several atomicities.

Mr. Davis's first attempt is worthy of all praise, but he himself will be the first to acknowledge that it is susceptible of improvement. We have privately heard that it has received the warm approbation of one of the most highly placed chemical teachers in this country, who has recommended it to his students, a piece of advice which we most cordially endorse. Metz, it is said, was taken by efficient *Kriegs Spieler*: we shall be glad to see some correspondingly strong chemical fortress reduced by Harnoezin players.

CORRESPONDENCE.

CLEMENTS' MANUAL OF ORGANIC CHEMISTRY.

To the Editor of the Chemical News.

SIR,—In a review of my book in your valuable journal of the 20th inst. several misstatements are made which I hope you will allow me to call attention to. Your reviewer states in the 31st line from foot of page 275 that no explanation is given of isomerism and polymerism. For isomerism please refer to pages 29, 47, 248, 254, 257, and 262, and for polymerism refer to page 262. Surely six or seven pages on this subject is sufficient in a work of less than 280 pages. In the line following it is stated that "no explanation is given of the difference between empirical and rational formulæ." Empirical formula is referred to on pages 3, 7, 9, 184, 192, 194, 196, 198, 202, and 223, and rational formula is referred to on pages 7, 8,

9, 10, 223, 229, and 254. I think in justice your reviewer should consider these references sufficient on this subject, considering that all students take up inorganic before organic chemistry, so that they would be well acquainted with the difference between rational and empirical formulæ. Moreover, page 223 alone should be sufficient to demonstrate to any student the essential difference between these formulæ. If the index is referred to, many of the other remarks will be found unnecessary, even including glycerin, &c. Furthermore, your reviewer states that "there is no evident distinction made between primary, secondary, and tertiary alcohols, nor between the different behaviour of aldehyds and ketons on oxidation." Pages 63 and 78 refer especially to the latter bodies, and the former is not borne out by the facts of the case. If your reviewer had thoroughly examined the book and referred to the index he might have omitted at least one-half of his review, which is both inaccurate and misleading.—I am, &c.,

HUGH CLEMENTS.

June 24, 1879.

Mr. Clements in his note says that "had the reviewer thoroughly examined the book and referred to the index he might have omitted at least one-half of his review." I beg to state that not only was the book thoroughly examined before the review was written, but after it was written the statements therein made were carefully compared with the index. I have looked up all the references he mentions in his letter, and find that, with two exceptions, they are quite irrelevant and frivolous. These two exceptions (pages 248 and 262), which refer to the explanation of isomerism and polymerism, though good explanations, are both given in the answers to the questions set by the Science and Art Department, and not in the *text* of the book itself, which is the point in question. In fact, the references which Mr. Clements gives are not to explanations of the terms in question, but merely references to the page where these terms are used; e.g., reference to page 47 for explanation of isomerism, "When it is distilled with CH_3HSO_4 methyl ether is formed, which is isomeric with ethyl alcohol." Also the reference to p. 184 for explanation of empirical formula—"What is the empirical formula of a substance possessing the following percentage composition:—C=40.0, H=6.66, O=53.34. Dividing by the atomic weights of C, H, and O respectively, we obtain 3.33 for C, 6.66 for H, and 3.33 for O, and dividing by the smallest we get 1, 2, 1 respectively for C, H, and O, i.e., the formula CH_2O . And the substance may therefore be $\text{C}_2\text{H}_4\text{O}_2$, acetic acid; $\text{C}_3\text{H}_6\text{O}_3$ lactic acid; $\text{C}_6\text{H}_{12}\text{O}_6$ grape-sugar." The references on pages 192, 194, 196, 198, 202, 223 are all of a kind exactly similar to the latter of the above two examples. The reference to page 254, as regards the explanation of rational formula, is "What is the rational formula of glycerine? $\text{C}_3\text{H}_5\text{H}_3\text{O}_3$." These are samples of the references which Mr. Clements gives for finding in his book explanations of rational and empirical formulæ, isomerism, &c. Mr. Clements does well to use in his letter the words "referred to" on such a page, for it is quite true the terms are there referred to, but this is no contradiction of the statements made in the review, where it is said that "No explanation is given of" the terms in question. He, in fact, *talks about* these terms, but gives no *explanation* of them. Page 223 is by no means an explanation, as Mr. Clements seems to think, of the terms rational and empirical formulæ, and even if it were it is not in the text of the book, but in the questions set by the Science and Art Department. With regard to glycerin, if Mr. Clements will again consult the review, he will find that the words "special reference" were used (and used advisedly) because though glycerin is often mentioned by name, yet no special statement is given (except in the case referred to in the review) of its general chemical reactions and relations to other compounds. On looking up the references on pages 63 and 78, as given by Mr. Clements,

I cannot find that any evident distinction whatever is made between the behaviour of aldehyds and ketons on oxidation. As regards the statement in the review that no evident distinction is made between primary, secondary, and tertiary alcohols, which Mr. Clements says "is not borne out by the facts of the case," how is it that he does not give a reference showing where the distinction may be found? I observe that on page 246, in the answers to the questions set by the Science and Art Department, an "attempted" explanation is given, but this is not all clear, and I believe that the average student would quite fail to understand it. It would seem that for some of the more important parts of organic chemistry we must consult not the "Manual" itself but the answers to the questions at the end of the book.

THE REVIEWER.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 24, June 16, 1879.

Observations on M. Lamansky's Paper entitled "On the Law of Stokes."—E. Becquerel.—The phenomena of fluorescence do not depend on a simple change in the refrangibility of the luminous rays which fall upon a body, but on a complete transformation of the vibratory movement. The incident rays excite vibrations in the substance, which then emits light whose composition does not seem simply connected with the nature of the incident vibrations.

Vapour-density of the Bihydrosulphate of Ammonia.—H. Sainte-Claire Deville.—The author gives the details of his determination of the vapour-density of neutral ammoniac sulph-hydrate, NH_4S , taken at the temperature 99.5° .

Electro-dilatation of the Coatings of Leyden Jars.—M. Duter.—The author has undertaken experiments which show that the cause of the expansion is not electric pressure, but that we have here to do with a new property of electricity.

Expansion of the Glass of Condensers while Charged.—M. Righi.—The phenomenon of Fontana, Govi, and Duter is due to a transverse expansion of the glass. The instantaneous expansion, due chiefly to the polarisation of the glass, must be distinguished from the persistent expansion not previously observed and due to a liberation of heat.

The Basic Hydrosulphates of Ammonia.—M. L. Troost.—The author has obtained several compounds of hydrosulphuric acid with ammonia. One of them at 0° occurs in ortho-rhombic crystals which act powerfully upon polarised light, upon which the ordinary crystals of bihydrosulphate have no action. A second solidifies at -8° , but may be maintained in a state of superfusion down to about -25° . The third has not been solidified at -55° . The author is engaged with their perfect separation and the determination of their vapour-tensions.

A New Native Manganese Sulphate and a New Variety of Ferrous Sulphate.—A. Carnot.—The new manganese sulphate, Mallardite, contains 7 molecules of water, and was obtained from the Lucky Boy Silver-mine, near the Butterfield Canon, Utah. The other mineral, luckite, is a manganiferous ferrous sulphate from the same deposits.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 1, 1879.

On Betulin.—N. Franchimont.—The author gives an account of the researches of H. Wigman. He assigns to betulin the formula $\text{C}_{36}\text{H}_{60}\text{O}_3$, but has not succeeded in ascertaining its nature, which, on the one hand, appears analogous to the terpenes, and, on the other, to certain aromatic hydrocarbons of high molecular weight.

On Lactucon.—N. Franchimont.—This compound, $\text{C}_{14}\text{H}_{24}\text{O}$, is obtained from *Lactua altissima*, and appears a homologue of camphor and of the zeorin of Paterno.

Acetate of Zinc.—N. Franchimont.—The crystalline acetate, contrary to the statements of most writers, contains only two molecules of water.

Colouring Matter of Santal and Calliatura Wood.—N. Franchimont.—The colouring matter of these dyewoods is identical, and may be represented by the formula $\text{C}_{17}\text{H}_{16}\text{O}_6$. Calliatura wood is the richer in this compound. The pure colour, on fusion with caustic potash, yielded acetic acid, resorcin, and probably proto-catechuic acid, and pyro-catechin.

Influence of the Chemical Composition of Water in the Preparation of Raw Silk.—Communicated to the Instituto Technico Superiore di Milan.—In silk distinguished the soluble constituents, "varnish," "gum" and colouring matters, and, on the other hand, the insoluble fibre. The soluble constituents give raw silk its brightness, colour, and strength, and should therefore be preserved as far as possible. For the purpose of unwinding the cocoons the natural gum should be softened, but not dissolved. According to Franceson, silk, if deprived of all its soluble constituents, loses at the same time its strength and elasticity. The authors find that though the loss of strength is proportional to the loss of soluble matter the elasticity is but slightly diminished. In order to minimise the loss sustained in softening the cocoons hard waters are used, and soft waters are artificially modified by the addition of sulphate of lime and carbonate of soda. Silks which are to be dyed bright colours, however, should be spun out of soft water.

Action of Mono- and Diphenylarsinic Acid upon the Animal Organism.—H. Schulz.—Both these substances are poisonous and agree in their action with ordinary arsenical compounds.

Action of Cacodylic Acid (Dimethylarsinic) upon the Animal Organism.—H. Schulz.—According to the experiments of Bunsen, Kürschner, and others, cacodylic acid has been regarded as not poisonous. The author, on careful experimentation with a pure sample freed from arsenic penta- and trioxide by repeated recrystallisation out of absolute alcohol, found it poisonous, the usual symptoms being produced in the subjects.

Action of Hydrocyanic Acid upon Epichlorhydrin.—J. v. Hörmann.—The product of the reaction is chlorohydroxybutyric acid.

Succinyl Compounds of the Toluydins.—G. v. Berzelius.—In this preliminary communication the author describes toluylyl-succinimid, $\text{C}_{11}\text{H}_{11}\text{O}_2\text{N}$, obtained by heating together equal mols. of succinic acid and ortho-toluydic acid.

Schützenberger's Acetates of Chlorine and Iodine.—B. Aronheim.—The author considers it doubtful whether these products are definite compounds or mere mixtures.

Action of Nitrous Acid upon Resorcin Ether.—B. Aronheim.—The result is an acid ethyl-ether of mono-nitro-resorcin.

On Digallic Acid.—H. Schiff.—A controversial paper, having reference to Freda's memoir (xi., 2033).

Certain Derivatives of the Hydroquinones.—R. Nietzki.—The author describes his researches on mono-nitro-diethyl-hydroquinon.

Reply to H. O. Hesse.—C. Rice.—A. controversial paper on the quinon-alkaloids (xi., 1549).

Certain Species of Sugar.—M. Hönig and M. Rosenfeld.—A preliminary communication on the sodium compounds of fructose and lactose.

On Para-phenylen-diamin.—A. Krause.—The author describes here the action of chloride of lime in solution upon para-phenylen-diamin hydrochlorate, the behaviour of aqueous hydrochloric acid with the chlorine product $C_6H_4Cl_2N_2$, bichlor-bibrom-quinon, and monochlor-monobrom-anilic acid.

Behaviour of Alkaline Solutions of Alumina with Sulphuretted Hydrogen.—G. Lösekann.—These solutions, on saturation with hydrogen sulphide, deposit all their alumina as hydrate, which, when washed, dissolves in acids without evolution of hydrogen sulphide. No alumina can be detected in the solution. Alkaline solutions of chrome behave in a similar manner.

Splitting-up of Bichlor-acrylic Acid by Alkalies.—O. Wallach and O. Bischof.—A detailed examination of the reaction by which chlor-acetylen is produced out of bichlor-acrylic acid.

Reactions of Silver Ultramarine.—K. Heumann.—Two atoms of the silver are combined in a different manner from the third and are more readily substituted by sodium. That atom of silver which on the decomposition of silver-ultramarine by acids is separated out in combination with sulphur is not removed by the action of sodium chloride, but retains its place in the molecule.

Relative Affinity of Oxygen for Hydrogen and Carbonic Oxide.—A. Horstmann.—If carbonic acid is added in varying proportions to mixtures of hydrogen and carbonic oxide with insufficient oxygen, less of the carbonic acid and more of the hydrogen will be burnt than in parallel experiments without carbonic acid.

Eikosylen, a Derivative of the Paraffin of Lignite.—E. Lippmann and J. Hawliczek.—Eikosylen is homologous with cetylen and must be regarded as a high member of the acetylen series.

On Amyliden-anilin.—E. Lippmann and W. Strecker.—This compound is the first instance of a new series of tertiary bases, where in an amine molecule the aldehyd residue takes the place of both H.

Nitrocuminol and its Derivatives.—E. Lippmann and W. Strecker.—An examination of nitrocuminol, nitrocuminic acid, and amido-cuminic acid.

Compounds of Cobalt and Nickel Chloride with Tar Bases.—E. Lippmann and G. Vortmann.—This paper does not admit of useful abstraction.

Researches on the Bessemer Process (III. The German Bessemer Process).—F. C. G. Müller.—In the German modification of the Bessemer process the initial temperature is 1400° , as against 1200° in the original English process. Manganese, if exceeding 2.5 per cent, has an unfavourable influence on the working of the German process, and all well-managed works endeavour to keep its quantity below 2 per cent.

Gases Occluded in Iron and Steel.—F. C. G. Müller.—The respective proportions of the gases present in Bessemer steel before and after the addition of the charge of spiegeleisen and in Martin steel are:—

H		88.8	77.0	67.8
N		10.5	22.9	30.8
CO		0.7	—	2.2

Further Communications on the Formation of Xanthoid Bodies from Albumen.—H. Krause and G. Salomon.—The authors find that only the incipient stages of putrefaction are favourable to the formation of hypoxanthin.

Oxidation-products of Cinchonin.—W. Königs.—The author has examined the products obtained by distilling

cinchonic acid with lime. Among the products obtained is oxy-cinchonic acid.

Constitution and Synthesis of Carbo-triphenyl-triamin.—W. Weith.—This synthesis is effected from aniline and para-nitrobenzoic acid.

Formation of Hydro-cinnamic Acid during Pancreatic Digestion.—E. and H. Salkowski.—Hydro-cinnamic (phenyl-propionic) acid is obtained during this process along with butyric and valerianic acids.

On Daphnetin.—Carl Stünkel.—The composition of pure daphnetin is $C_9H_6O_4$. Among its derivatives mono-acetyl-daphnetin, tetra-brom-mono-acetyl-daphnetin, and mono-benzoyl-daphnetin are here analysed and described.

Ethers of Tribasic Formic Acid.—A. Deutsch.—An examination of ortho-formic ethyl-ether, with the corresponding methyl-propyl, iso-butyl, and iso-amyl compounds.

Contributions to the Volumic and Steric Laws. H. Schröder.—This memoir gives the steres of the sodium, lithium, and calcium compounds.

MISCELLANEOUS.

Non-poisonous Character of Copper.—M. Galippe, in a paper read before the Biological Society of Paris, described the following experiment:—A rabbit received daily for six months 2 grammes (?) copper acetate. At the end of this time it was served up at the table of the learned chemist. The liver weighed 70 grms., and contained 13 centigrams of copper. M. Galippe partook of it, and has suffered no inconvenience.

Chemical Society Research Fund.—The following grants have just been made from the Research Fund of the Chemical Society:—

£30 to Mr. W. Whiteley Williams, for experiments on an Improved Method of Organic Analysis.

£25 to Mr. M. M. Pattison Muir, of Caius College, Cambridge, for Determining the Physical Constants and Chemical Habitudes of certain Bismuth Compounds.

£15 to Mr. J. M. Thomson, for Experiments on the Action of Isomorphous Bodies in Exciting the Crystallisation of Supersaturated Solutions.

£50 to Dr. Wright, for continuing his Researches on Chemical Dynamics.

£25 to Mr. F. D. Brown, for continuing his Researches on the Theory of Fractional Distillation.

£30 to Mr. Bolas for an Investigation of certain Chromium Compounds.

£20 to Mr. F. R. Japp for an Investigation of the Action of Organo-zinc Compounds on Quinons.

£100 (the De la Rue donation) to Dr. H. E. Armstrong, for the Determination of certain Physical Properties especially Refraction Indices, of Typical Chemical Compounds.

The Treasurer, Dr. Russell, informs us that Dr. De la Rue has announced his intention of presenting the fund with £100; this will be the third donation of that amount which the fund has received from him.

NOTES AND QUERIES.

Tetrachloride of Carbon.—Will any of your readers kindly state the simplest and most practical mode of producing tetrachloride of carbon?—CIC.

Combustion of Magnesium.—If a small bundle of magnesium ribbon is burnt in a porcelain crucible, incomplete combustion ensues. If the ignited mass is then treated with HCl, H is evolved, and a series of small explosions takes place, the H taking fire. A fishy smell is at the same time evolved, rather like trimethylamin. Is the presence of a nitride sufficient to account for these phenomena? The porcelain crucible is slightly attacked, and a black insoluble stain left, which according to Gmelin is due to silicon.—S.

ERRATUM.—P. 260, col. 1, line 15 from top, for phosphoric read propionic.

A Chemist (21), who has had practical experience in works, desires Re-engagement. Good references; no objection to go abroad.—For particulars, &c., address, X. T., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

A Chemist, who has had several years practical experience, is open to an Engagement. During the last six years has been engaged in the Alkali Manufacture.—Address, W. H. R., Fir Cottage, Weston, Runcorn.

Chemist, who has studied under the first Professors in England and Germany, and has had many years experience in theoretical and practical chemistry, and been in works, desires Re-engagement in Works or Laboratory, preferably in Brewery. Very highest testimonials and references.—Address, "Eosine," 3, Wimpole Street, Cavendish Square, W.

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A French Gentleman, speaking English fluently, thoroughly conversant with English and French pharmacy, and an Associate of the Pharmaceutical Society, wishes a Re-engagement; London preferred. First class references, &c.—Address, Z., Laboratory, 143, New Bond Street, W.

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Wanted, a German Chemist who has a practical knowledge of acetate soda making, and of the separation of arsenic from sulphuric acid.—"Civis," CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

Wanted, by a Young Man (29), a Situation either in a Works or Town Laboratory. Ten years experience and accustomed to management of men. Has had considerable practice in water, manure, sugar, and general analyses; also in original investigations. Good references.—Address, F.C.S., Iron-deg, Abergyle.

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THE CHEMICAL NEWS.

VOL. XL. NO. 1024.

A NEW APPLICATION OF RAPID OXIDATION BY WHICH SULPHIDES ARE UTILISED FOR FUEL.

By JOHN HOLLWAY.

[MR. JOHN HOLLWAY has forwarded to us for publication the following account of his new application of rapid oxidation by which sulphides are utilised as fuel. Many who may not have read the longer account given in Mr. Hollway's paper read before the Society of Arts, February 12th, 1879, will probably be glad of this summarised account of the process.]

This process has for its object the utilisation of the heat generated by the rapid oxidation of certain mineral substances which have not hitherto been used as sources of heat for smelting operations. The heat thus obtained is employed in the reduction of the furnace charge, which may be composed partly of sulphides and partly of silicious ores. A current of air is forced through molten sulphides, by which means they are very rapidly oxidised. Great heat is thus developed, rendering the process of smelting a self-supporting operation; therefore no extraneous fuel is required, excepting that employed in raising steam for the blowing engines; where, however, water power is available, steam can be dispensed with, in which case all the carbonaceous fuel necessary for the operation is a little coke to start the furnaces, which stands in the same relative position to the ores as wood does to coal in the lighting of an ordinary fire.

It is well known that pyritous minerals are readily combustible, but the best means of utilising the heat-producing property of metallic sulphides is not so apparent as would at first sight appear. Of these sulphides only iron pyrites is sufficiently combustible at a low temperature to burn in the open air, the mass being raised to the temperature at which the oxidation takes place solely by the union of sulphur and iron with atmospheric oxygen. In Spain there are numerous deposits of poor cupreous pyrites, and the Rio Tinto and Tharsis Companies annually treat at their mines about one million tons for the extraction of copper only, which does not average 2 per cent. The process employed consists essentially in roasting the pyrites in heaps in the open air, dissolving out the copper from the roasted material, and precipitating it from the solution by means of iron. These operations extend over several months, any gold or silver contained in the ore is lost, and the iron and sulphur are also wasted. The sulphur passes into the air as an obnoxious and annoying gas, desolating the country for miles around the works.

From the earliest ages, carbon has been considered a necessity in all metallurgical operations. The first reduction of metals by means of carbon forms a connecting link between the age of stone and the commencement of civilised art. It is well known that carbon burns at widely varying temperatures, as, for example, in our bodies, in a common coal fire, or in a furnace. A great deal of thought has been devoted to the subject of economising carbonaceous fuel, and great advances have been made in this direction; yet the expenditure of coal or coke necessary, say to melt a given quantity of metal, still far exceeds the theoretical limit. The main causes of this discrepancy may be accounted for as follows:—

1st. Only part of the oxygen of the air passing into a furnace acts on the material to be burnt.

2nd. The oxygen is not brought in contact with the combustible matter with sufficient rapidity to obtain the necessary temperature for the operation.

3rd. Gases pass off hot and unburnt. These are now, however, frequently utilised.

There is one metallurgical operation in which the first two sources of loss are avoided, viz., "Bessemer's," where, by blowing air through molten crude iron, a very high temperature is attained by the combustion of small quantities of carbon and silicon contained in the crude iron; this is, however, not the case in the process of puddling, where the oxidation is spread over a considerable period of time, although the same constituents are frequently burnt in similar proportions. But even in the Bessemer process the carbon is only half burned, and a large amount of heat escapes with the carbonic oxide and nitrogen.

When, however, thin streams of air are forced through molten sulphide of iron lying on a tuyère hearth, a high temperature is produced by the perfect combustion which ensues in the midst of the sulphides, and no unburnt gases, excepting nitrogen and sulphur vapour, escape from the surface of the molten mass. The hot nitrogen and sulphurous acid may be caused to act upon iron pyrites and other mineral matter, and when pyrites is thus heated an atom of sulphur held in feeble combination is in great part expelled, and thus is obtained molten protosulphide of iron, which is subsequently burnt by the oxygen of the air driven in at the lower part of the furnace, thereby producing the heat necessary for continuing the operation. The process may be defined as a system of fractional oxidation, in which the numerous constituents of a complex furnace charge can be separated from each other and concentrated in different parts of the apparatus, the heat necessary for the operation being obtained by the combustion of a portion of the less valuable constituents.

The principal ores of all our ordinary heavy metals, except manganese and tin, are sulphides. Iron, although largely occurring in an oxidised form, is abundantly found in combination with sulphur; and bi-sulphide of iron, or iron pyrites, is an example of sulphurous and combustible minerals associated with the iron and sulphur; in iron pyrites are invariably found small quantities of other metals, notably cobalt, nickel, copper, silver, gold, lead, zinc, and arsenic. Of these zinc is almost as combustible as iron itself, while lead and arsenic readily volatilise as sulphides, and cobalt, nickel, and copper are distinctly less readily oxidisable than iron, while silver and gold do not oxidise under these conditions; hence, in supplying air to such material, the iron is the first of the elements to suffer oxidation, so that if the oxidation be arrested before the whole of the iron has been burnt, the cobalt, nickel, copper, silver and gold present will be found in the unburnt portion. This principle finds a parallel in the Bessemer process of treating pig-iron for the manufacture of steel, where a current of air is caused to bubble up through a bath of molten crude iron, the silicon is first oxidised, and is closely followed and to a great extent accompanied by the carbon, and no large amount of iron suffers oxidation until the whole of the silicon and carbon have been burnt out of the molten material.

The experiments made at Messrs. Cammell's works, at Penistone, in a Bessemer converter, have proved that by blowing air through molten sulphide of iron, the iron and a portion of the sulphur are oxidised, and if the oxidation is arrested before the combustion of the iron is complete a heavy matt or regulus is obtained, which contains but a small proportion of the iron of the ore, but practically the whole of the greater part of the copper and other less oxidisable metals. In one of these experiments the molten sulphides were run into the converter from a cupola, in which they had been previously melted, and the temperature was kept up until the operation was discontinued—viz., for a period of ten hours, without the use of any carbonaceous fuel, the heat being entirely derived from the oxidation of the iron and a portion of the sulphur of the lumps of pyrites, which were continuously thrown into the mouth of the converter. A Bessemer co-

verter being unsuited for the collection of the gaseous products, the later experiments have been made in a series of cupola furnaces belonging to Messrs. John Brown and Company, Limited. These experiments have proved the possibility of obtaining a valuable regulus, a slag nearly free from copper, and a considerable quantity of crude sulphur. M. Pourcel, the well-known chemist of the Terrenoire Company, has also made some very interesting experiments, having treated by this method a cupriferous sulphide of antimony containing lead and zinc, using heavy spar and silica as fluxes; he obtained a regulus containing the whole of the copper in the form of sulphide, a slag of light specific gravity, and the lead, zinc, and antimony as two separate sublimes which were condensed in different parts of the apparatus, owing to the superior volatility of sulphide of lead over the oxides of antimony and zinc. In the experiments at Penistone and at Sheffield a cold blast of air was employed, and the gases which passed from the converter or furnace into the open air carried away with them a large amount of heat. In practice, however, it would be economical to employ a hot blast, which could be heated by the waste heat from the escaping gases. It is remarkable that the least valuable metals—viz., iron and zinc—generate by their combustion the largest quantities of heat.

The process may be employed for the reduction of even the more volatile metals; for example—Mr. A. H. Allen, of Sheffield, has thus obtained metallic antimony simply by the oxidation of sulphide of antimony. It is well known that sulphide of lead reacts upon oxide of lead with the production of metallic lead and sulphurous acid. If, therefore, a limited amount of air is blown into molten sulphide of lead, the oxide thus formed in the lower part of the furnace will, in passing upward, come in contact with the hot sulphide of lead, and metallic lead will result with the evolution of sulphurous acid. The furnace having a quiescent hearth below the tuyères, the metallic lead will collect there, and can be from time to time withdrawn. A limited amount of air must be employed, because if it is driven in too quickly, the sulphide of lead would rapidly distil off. In thus treating argentiferous lead ores, the silver (and gold if present) would be found with the first metallic lead reduced. When thus treating galena the furnace should have a basic lining.

The process is peculiarly suitable:—

1st. For the treatment of metalliferous substances which cannot be advantageously treated by other processes. For the extraction of sulphur by distillation, and simultaneously for the concentration and separation of cobalt, nickel, copper, silver, and gold from minerals in the form of metallic regulus, while lead, zinc, antimony, arsenic, &c., accrue in the sublimes.

2nd. For the treatment of complex ores, for example—Grey antimonial copper ores, such as those experimented on by M. Pourcel. Ores similar to those worked at the well-known Bottino Mines, Seravezza, in the Italian Apennines, which contain thirteen or fourteen heavy metals, including silver and lead, for which latter alone they have been worked for centuries. The blende of lead mines in Derbyshire termed "muck," usually thrown away by the miners, because the large quantity of lead with which it is associated renders the zinc obtained from it worthless.

3rd. For the treatment of auriferous and argentiferous pyrites. It is well known that in practice it is not possible to obtain the whole of the gold from pyrites by amalgamation with quicksilver, because the presence of sulphur and arsenic sickens and flours the mercury, whereas by fusion the whole of the silver and gold present is obtained.

4th. For the treatment of pyrites containing even only small percentages of cobalt, nickel, and copper, which are thus concentrated into a rich regulus, whereas this result is now only obtained by very tedious processes of alternate roasting and reduction. Such ores containing 10 per cent and even 12 per cent of copper exist in South Ame-

rica and many other parts of the world, but are not at present capable of economic treatment, owing to the difficulty of obtaining a sufficient supply of cheap fuel. The process can also be advantageously applied to the treatment of richer ores of copper such as are at present smelted at Swansea.

5th. For the treatment of poor lead ores. If such ores are added to a furnace charge of cupreous pyrites, the silica they contain will be utilised and combine with the resulting oxide of iron to form slag, the galena will be volatilised and be recovered as a sublimate, while any silver present will enrich the regulus. At present, by a costly process of crushing and washing these ores, the galena is concentrated, although a large proportion is left with the *débris*, and passes with the water into the streams, rendering the existence of fish in such waters impossible. The water power now used for washing the ore could, in many cases, be employed for producing the blast.

When thus treating cupriferous iron pyrites, four products are obtained:—

1st. A matt or regulus containing from 30 to 50 per cent of copper, any traces of cobalt, nickel, silver, or gold the ore may contain, the rest of it being iron and sulphur it has a specific gravity of $4\frac{1}{2}$ to 5.

2nd. A slag consisting of silicate of iron from the resulting oxide of iron combined with the silicious matter contained in the ore and the fluxes added.

3rd. Sublimed sulphur, more or less mixed with volatile compounds of lead, zinc, and arsenic.

4th. Sulphurous gases, consisting mainly of sulphuro acid and nitrogen.

The regulus closely resembles, and will replace, coarse metal of the Swansea copper process, which is only obtained at considerable cost of labour, time, and carbonaceous fuel. When, however, sulphides of iron and copper present in the bath are treated continuously by a blast of air a point is at length arrived at when the whole of the iron is oxidised, and the regulus in the bath consists of sub sulphide of copper. If, now, a limited supply of air is introduced, the copper is reduced to the metallic state with the evolution of sulphurous acid.

The slag obtained in the Penistone experiments was essentially silicate of iron containing about 50 per cent of iron and 29 per cent of silica. It had a density of about 3.8 to 4, and was in composition somewhat allied to the copper-smelters' ore furnace slag and to the tap-copper of the iron-puddler. By the addition of calcareous materials the specific gravity of the slag is so reduced as to cause it to separate readily from the regulus which collects below it. In one of the later experiments, when lime was used, the proportion of copper lost in the slag was very small. This is, of course, a most important point, for when dealing with ores containing but little copper, the presence of even a small percentage in the slag means the loss of a considerable proportion of the copper present. These slags can be utilised for the manufacture of steel, being practically silicious iron ore free from phosphorus, and their reduction in a blast-furnace can be profitably effected, as the proportion of iron present is high as compared with the weight of the material; indeed, it may be possible to reduce them while in a molten state.

By re-subliming the crude sulphur, it can be freed from all impurities except arsenic, and at the works of Messrs. John Hutchinson and Co., Widnes, this is eliminated by means of polysulphide of calcium.

As a certain proportion of the sulphur of the minerals suffers combustion, the resulting sulphurous gases contain from 14 to 15 per cent of sulphurous acid, and hence the proportion of sulphurous acid to nitrogen is nearly identical with that of the gases produced by roasting pyrites in the kilns employed by vitriol manufacturers, and can therefore be used with equal advantage for the production of vitriol in leaden chambers. This appears to be the simplest solution of the great problem how to smelt copper without

causing a nuisance to the surrounding neighbourhood, although a similar result might be obtained by collecting and liquefying the sulphurous acid.

The more incombustible materials it is found practicable to employ without too great a loss of temperature, the wider will become the application for the process; for there are many ores, including silicates and carbonates, containing metals in the form of oxides, which might be conveniently smelted by mixing them with a sufficient proportion of pyritous ores to effect their reduction; in fact, one of the chief practical questions connected with this process is how far it may be trusted to effect the smelting of ores or furnace-charges containing comparatively moderate proportions of sulphides. It is evident that it will almost entirely obviate the necessity for using carbonaceous fuel, at least as far as the production of a regulus is concerned, and consequently the localities in which smelting operations may be advantageously carried on are thus greatly multiplied. One of its chief merits is that it is equally applicable, with comparatively little extra cost in the working, to very poor and very rich ores; for however small the resulting regulus, it will contain nearly the whole of the cobalt, nickel, copper, silver, and gold present in the furnace-charge, while any lead, zinc, antimony, and arsenic will be obtained as sublimes.

RESEARCHES IN CHEMICAL EQUIVALENCE.*

PART III.† NICKELOUS AND COBALTOUS SULPHATES.

By EDMUND J. MILLS, D.Sc., F.R.S., and J. J. SMITH.

ALTHOUGH the chemistry of nickel and cobalt is interesting from many points of view, it is more especially attractive from the probable isomerism of these metals. Their combining proportions, in fact, according to the most valuable evidence we possess,‡ appear to be entirely the same. We, therefore, thought it very advisable to inquire on what terms they might prove to be mutually equivalent; and the particular equivalence we have examined has been equivalent precipitability of the sulphates, by sodic hydrate, from an aqueous solution.

I. Preparation of the Salts.

The pure cobaltous salt was prepared by converting some excellent commercial crystallised chloride into luteo-chloride, the process employed having been already described by one of us.§ The luteo-chloride was purified by precipitation with hydric chloride, and the mixture of oxides it left behind on ignition was evaporated with redistilled hydric sulphate.

The pure nickelous salt was prepared from a sample of nickelous chloride which contained copper, lime, and iron, but no cobalt. The copper present was precipitated with hydric sulphide, and the nickel in the filtrate was precipitated by hydric oxalate, in an acid solution. The nickelous oxalate was washed thoroughly with dilute hydric nitrate, ignited, and the oxide so formed heated with pure hydric sulphate, and so converted into sulphate.

II. Method of Separating Nickel from Cobalt.

We had next to select a method for the quantitative separation of nickel from cobalt. The first to which we had recourse was Liebig's,|| which consists in adding hydric cyanide and potash to the mixed saline solutions, thereby forming nickelopotassic cyanide and potassic cobalticyanide: these new compounds are boiled with

freshly precipitated mercuric oxide, which throws down the nickel as oxide and cyanide. The cobalt in the filtrate is then precipitated as mercurous cobalticyanide. On the ignition of these precipitates, nickelous oxide and cobaltic oxide are respectively left behind. We converted the oxides into sulphates and weighed them as such. (Throughout the whole of these experiments the nickel and cobalt were always weighed as sulphates.) We found, however, that after the expulsion, by ignition, of the mercury in the precipitates, the oxides were left in so dense and compact a state that it was only with the greatest difficulty that they could be converted into sulphates. To remedy this a weighed quantity of pure baric sulphate was added to the liquid before precipitation; the precipitate, becoming mixed with the baric sulphate, was thus spread over a large surface, so that the oxide was obtained in a finely divided state, and easily converted into sulphate.

In the experiments with this process, the nickel was invariably found too high. This was at first thought to be due to alkali which might cling to the precipitated oxide; and, on testing, some potash was actually found in the precipitate, which had been thoroughly washed. To remove this source of error the nickelous sulphate, instead of being weighed as such, was dissolved in water, excess of ammonia added, and the nickel deposited on a platinum crucible by means of two Grove's cells.

But the nickel was, with this method, still found too high, and we then thought that the separation had not been perfect. On testing the deposit, a considerable amount of cobalt was, in fact, found in it, and the operation required to be repeated three times before the separation was complete.

The next process we tried was devised by Rose.* It consists in saturating with chlorine a very dilute solution of the mixed salts, adding baric carbonate in excess, and allowing to stand from twelve to eighteen hours, with occasional shaking. The cobalt then falls as sesquioxide, the nickel remaining in solution.

But the results obtained by this method, as was also the case with Henry's modification of it (which consists in substituting bromine for chlorine) were very variable. In the former case the cobalt was generally found too low, even after standing over eighteen hours; the result appearing to depend a good deal on the liquid being shaken up at regular intervals, which cannot be very conveniently done during an entire period of eighteen hours. In the latter case the cobalt was generally found too high, nickel being precipitated along with it.

The last method tried, and the one finally adopted, is due to Gibbs.† In this the neutral solution of the sulphates is boiled with plumbic peroxide. The cobalt is then precipitated as a higher oxide, while the nickel remains in solution, along with a small quantity of lead. (The author of this process does not claim very great accuracy for it, but we have found it to be of adequate accuracy.

The manner in which we operated was as follows;—The perfectly neutral solution of the mixed sulphates was boiled for half an hour with plumbic peroxide, about 7 grms. of the peroxide being taken to 1 grm. of cobaltous sulphate, and the volume of the liquid being about 100 cub. centims. The liquid was then filtered, and the filtrate evaporated to about 20 cub. centims. Some aqueous hydric sulphide was then added, and the small quantity of plumbic sulphide formed filtered off. The solution of nickelous sulphate was then evaporated to dryness in a weighed crucible, ignited, treated with a little hydric sulphate, heated to very dull redness, and weighed. The precipitate containing the cobalt, with excess of plumbic peroxide and some sulphate, was boiled with hydric chloride until dissolved. Water was then added, and the lead precipitated with hydric sulphate and alcohol. The filtrate from the plumbic sulphate was then evaporated to dryness, heated till the excess of hydric sulphate was

* A Paper read before the Royal Society, June 19, 1879.

† For Part II. see *Proceedings of the Royal Society*, vol. xxviii., p. 270.

‡ Russell, *Journal of the Chemical Society* (1869), p. 294.

§ *Philosophical Magazine* (4), xxxv., p. 245.

|| *Ann. Ch. Pharm.*, lxx., 244.

* *Pogg. Ann.*, lxxi., 545.

† *Sill. Am. J.*, xiv., 203.

Nickelous salt taken.	Cobaltous salt taken.	Cobaltous salt taken (corr'd.)	Total precipitate.	Nickelous salt precipitated.	Cobaltous salt precipitated.	Cobaltous salt precipitated (corr'd.)	Sodic sulphate precipitated.	Temperatures.	Number of experiment.
gram.	gram.	gram.	gram.	gram.	gram.	gram.	gram.	Degrees.	
0.1	0.9	0.8093	0.9155	0.1050	0.8105	0.7198	0.0105	7-8	I.
0.2	0.8	0.7581	0.8667	0.1980	0.6687	0.6268	0.0110	6-7	II.
0.3	0.7	0.6268	0.8980	0.2765	0.6215	0.5483	0.0135	7-8	III.
0.4	0.6	0.5598	0.8650	0.3510	0.5140	0.4738	0.0105	7-8	IV.
0.5	0.5	0.4363	0.8885	0.4465	0.4420	0.3783	0.0090	7-8	V.
0.6	0.4	0.3613	0.8635	0.5295	0.3340	0.2953	0.0150	7-7.5	VI.
0.7	0.3	0.2808	0.8440	0.5825	0.2615	0.2423	0.0110	7-7.5	VII.
0.8	0.2	0.1408	0.8840	0.7080	0.1760	0.1168	0.0130	8-8.5	VIII.
0.9	0.1	0.0483	0.8765	0.7865	0.0900	0.0383	0.0105	7-7.5	IX.

driven off, and dissolved in water. A little hydric sulphide was then added, to remove a small quantity of lead still present, the filtrate evaporated to dryness, and the cobaltous sulphate weighed. The plumbic peroxide must, of course, be perfectly pure, because any impurity in it finds its way into the cobalt, and consequently makes that result too high.

In an experiment with this process 0.2500 gram. of cobaltous sulphate and 0.2500 gram. nickelous sulphate were taken, and the quantities found after the separation were 0.2505 gram. cobaltous sulphate and 0.2490 gram. nickelous sulphate. The estimation of the cobalt is rather a troublesome and tedious process; but when the mixture of nickel and cobalt can be weighed, and the nickel in it estimated, the cobalt being taken by difference, this is an admirable method to employ. The following are some results of the estimation of the nickel in a mixture of 0.2500 gram. of cobaltous sulphate and 0.2500 gram. of nickelous sulphate:—

	Nickelous sulphate taken.	Found.
(1.)	 0.2500	0.2480
(2.)	 "	0.2505
(3.)	 "	0.2490
(4.)	 "	0.2495
(5.)	 "	0.2480

Mean 0.2490

Probable error of a single determination 0.16 per cent.

III. Experiments on Equivalence.

Having then fixed on the method of separation, 1 per cent solution of nickelous and cobaltous sulphates were prepared, and a solution of sodic hydrate, of which 10 cub. centims. were capable of precipitating 0.8248 gram. of nickelous or cobaltous sulphate. This sodic hydrate was made from sodium, and kept in glass bottles coated internally with a thick layer of paraffin.

A series of nine experiments was made, in which the relative weights of nickelous or cobaltous sulphate present varied from 0.1 to 0.9 gram.; the total weight of nickelous and cobaltous salt, and the volume of the solution being, however, always the same—viz., 1 gram. and 100 cub. centims. The experiments were conducted as follows:—The bottles containing the solutions of the sulphates and the sodic hydrate were immersed in a trough into which there was a constant flow of water to bring them to a constant temperature. The necessary quantities of nickelous and cobaltous solutions were then carefully measured out, mixed, and the temperature observed. 10 cub. centims. of sodic hydrate were then added, the solution stirred vigorously, and the temperature again observed. The precipitate was then filtered off as quickly as possible (an aspirator being used to facilitate the filtration), and washed, first with cold and then with hot water. Three days' intermittent washing was required to free the precipitate from the undecomposed nickelous and cobaltous salts, cobaltous hydrate having, as is well known, a powerful attraction for cobaltous sulphate, thereby forming a basic salt. After washing, the precipitate was converted into sulphate by treatment with hydric sulphate, and weighed. The mixed sulphates were then dissolved in

water, and separated by Gibbs's method, the nickelous sulphate being weighed, and the cobaltous sulphate obtained by difference. It was found that a small quantity of sodic sulphate was always present in the precipitate, the washing having failed to remove it, and this required to be estimated and deducted. For this purpose the nickelous sulphate, after being weighed, was dissolved in water, and the nickel precipitated with baric hydrate. The barium in the filtrate was then removed with hydric sulphate, and the filtrate containing the sodic sulphate evaporated to dryness and weighed. The results obtained are comprised in the table at top of page.

In this table the precipitates are all returned as sulphates. The total possible amount of normal sulphate attainable with the constant quantity of sodic hydrate employed having been 0.8248 gram., we are able to calculate the entire composition of the precipitate thus:—From its total weight, the amount of sodic sulphate is first subtracted; from the residue the constant quantity 0.8248; the remainder, which is the cobaltous sulphate carried down with the hydrate, is deducted as a correction from the cobaltous sulphate originally taken and that which was precipitated, it having no share in the reaction we had to examine. Under "temperature" we give the temperatures of the reagents before and after mixture. The weight of precipitated sodic sulphate varies but little throughout the experiments, its mean value being 0.0115 gram. In a special determination, we precipitated, with the usual amount of sodic hydrate, 1.000 gram. of nickelous sulphate alone: after three days' washing, it was found to contain 0.0115 gram. of sodic sulphate. Hence we infer that when the two sulphates are present together, nickelous hydrate carries down sodic sulphate, cobaltous hydrate carrying down cobaltous sulphate.

IV. Discussion.

In the following discussion it will be understood that we refer to the tabular results already given.

If n represent a weight of nickelous sulphate taken, and v be the hydrate (calculated to sulphate) obtained from it through precipitation, then we consider ϕ in the expression $n = \phi v$ to represent the *precipitability* of nickelous sulphate: and similarly in $c = \phi \gamma$, ϕ represents the precipitability of cobaltous sulphate. In examining the numbers obtained with the nickelous salt, the best expression we could find for ϕ was $\phi = (a + \beta n)$. We first calculated the values of a and βn from all the determinations, and by the method of least squares, thus obtaining—

$$\phi = 0.98891 + 0.22571n.$$

It is, however, clear that a cannot be less than unity; moreover, the first weight of nickelous precipitate is somewhat higher than is possible, and we have thought well to reject it. With these amendments we finally obtain—

$$\phi = 1 + 0.21940n,$$

with a probable error of 0.02558 for a single determination, or 0.00904 for eight determinations. Hence we infer that the *precipitability of nickelous sulphate is directly proportional to its mass.*

In the case of cobaltous sulphate, on the other hand,

no such law holds good. After a very careful examination of the numbers, we could not find any satisfactory evidence of a change in precipitability with its mass, and consequently represent ϕ as a constant. The mean value of ϕ is in this case 1.1845, with a probable error of 0.02792 on a single determination, or 0.00931 for nine determinations. Thus ϕ is about equally well ascertained in both cases.

Our two equations may now be written—

$$\begin{aligned} n &= (+0.21940n)\nu \\ c &= 1.1845\nu \end{aligned}$$

In order to calculate in what proportions the two sulphates are equally precipitable, we have—

$$1 + 0.21940n = 1.1845,$$

whence $n = 0.84093$. The two sulphates, then, are equally precipitable when the weight taken of each is 0.84093 grm.

To calculate in what proportions the two sulphates must be mixed, to give equal weights of precipitate, we have—

$$\begin{aligned} \frac{n}{c} &= \frac{(1 + 0.21940n)\nu}{1.1845\nu} \\ n &= 1 - c \end{aligned}$$

If we put $\frac{\nu}{\gamma} = 1$, and combine these equations, the result is a quadratic, one of whose solutions is $c = 0.5168$, and consequently $n = 0.4832$. Using these values in the primitive equations, the results are, $\gamma = 0.4363$, $\nu = 0.4369$, which we may regard as substantially equal.

A highly important chemical relation is disclosed when the two sulphates are so conditioned as to be equally precipitable. We have seen that this is the case when $n = c = 0.84093$. Now the reciprocal of 0.84093 is 1.1892, a number differing by only the small amount of 0.40 per cent from 1.1845, or $\phi\gamma$. Deduced, as these values both are, from a series of experiments, we cannot hold their connection to be accidental. It may be thus expressed in symbols—

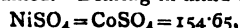
$$n = \frac{1}{\phi\gamma} = \frac{\gamma}{c}$$

or we may state the relation thus:—For an equal weight, nickelous and cobaltous sulphates are equally precipitable; the attraction of the one towards the reagent being then inverse to that of the other.

In order to ascertain whether this function admits of extension to other weights than 0.84093 grm. of the sulphates, we doubled all the masses in one of our previous experiments (V. in the table), and determined the chemical effect. The results were—

Uncorrected total precipitate	1.7840 grm.
Containing of nickelous sulphate	0.8925 "
The halves of these are	0.8920 "
	0.4463 "
Figures nearly identical with	0.8975 "
	0.4465 "

derived from Experiment V. By doubling the mass, we thus produce double the chemical effect. It is probable that this law is general; and, therefore, we infer that the reciprocal function we have noticed may apply to multiples of the weights to which, in our particular case, it specially appertained. Bearing in mind that—



we accordingly write the function thus—

$$\phi(\text{NiSO}_4) = \phi^{-1}(\text{CoSO}_4).$$

If we imagine some reaction—such, for example, as the combination of colouring matter with a tissue—influenced in one set of experiments by nickelous sulphate, and in an altogether different set by an equal weight of cobaltous sulphate, we can hardly conceive any ground for the development of a reciprocal function, such as we have experimentally

traced. On the other hand, it seems reasonable to suppose that when two bodies are simultaneously confronted with a single reagent, they both contend for its effect. Thus a chemical antagonism may arise between them by virtue merely of their being together; and thence the reciprocal function. So far as we are aware, the only other chemical function of the kind hitherto investigated is to be found in Chizynski's* examination of the partial precipitation, by ammoniac phosphate, of mixed calcic and magnesian chlorides. That chemist arrived at the conclusion, for which we consider his evidence to be adequate, that "equal masses of calcic and magnesian chlorides have always equal, but oppositely active, coefficients of affinity."

ON THE TOTAL ESTIMATION OF NITROGEN BY COMBUSTION, INCLUDING THE NITRO-COMPOUNDS.

By JOHN RUFFLE, M.R.A.C., &c.

In a previous note on this subject (CHEMICAL NEWS, vol. xxv., page 189) I stated that by modifying the well-known soda-lime process for a nitrogen determination by combustion, as much as 80 per cent of the nitrogen of a nitrate could be determined whether present in conjunction with ammoniacal salts or otherwise, and that under certain circumstances as much as 94 per cent of the nitrogen of a nitrate had been converted into ammonia in such combustion process. Also in the note I recognised the value of a ready method for the complete determination of the total nitrogen in a substance. During the seven years which have elapsed since the appearance of my note a great part of my leisure time has been spent in conducting experiments on this subject, and I at length have the great pleasure of reporting the following satisfactory results:—

Substance taken.	Percentage of Nitrogen by Theory.	Percentage found.
Soda nitrate— $\text{Na}_2\text{ON}_2\text{O}_5$	16.47	16.43
" " " " " " " "	"	16.32
" " " " " " " "	"	16.26
" " " " " " " "	"	16.56
Potash nitrate— $\text{K}_2\text{ON}_2\text{O}_5$	13.90	13.94
Lead nitrate— $\text{Pb}_2\text{ON}_2\text{O}_5$	8.45	8.21
Am. nitrate— $(\text{NH}_3)_2\text{N}_2\text{O}_5\text{H}_2\text{O}$..	35.00	35.04
Quin. slph. $2\text{C}_{20}\text{H}_{24}\text{N}_2\text{O}_2\text{H}_2\text{SO}_4$ } 7 aqua	6.41	6.30
Common pyroxiline used for col- lodion— $\text{C}_{18}\text{H}_{22}\text{O}_{15}(\text{NO}_2)_7$ (?)	(?)	11.64
" " " " " " " "	(?)	11.41
Powder from a Snider cartridge containing 75 per cent potash nitrate (?)	10.35 (?)	10.10
" " " " " " " "	"	10.12
An artificial manure containing superphosphate and nitrate soda 25 per cent = nitrogen— 4.11; sulphate ammonia 25 per cent = nitrogen—5.30 ..	9.41	9.28
" " " " " " " "	"	9.33
An artificial manure containing superphosphate and nitrate soda 25 per cent = nitrogen ..	4.11	4.24
" " " " " " " "	"	4.23
An artificial manure containing superphosphate and nitrate of ammonia 25 per cent = nitrogen	8.75	8.81
" " " " " " " "	"	8.92
A mixture containing, per cent— Suprphosphate. 70 Hair 10 = nitrogen 1.46 Sulph. amm. 10 = " 2.12 Nitrate soda 10 = " 1.64	5.22	4.99

* Ann. Ch. Pharm., Supp. iv., 226 to 233.

It will be seen that the above results give the nitrogen when present as a simple nitrate, or as a lower oxide of nitrogen, also when the nitrate is mixed in a complex substance like an artificial manure or gunpowder, and whether with or without the presence of organic matter. From these and other results the process employed will, I believe, through conversion of the nitrogen into ammonia by combustion, enable the ready and complete estimation of the total nitrogen in a compound or mixture in whatever form the nitrogen may be present—i.e., as an albuminous compound, ammoniacal salt, or as an oxide of nitrogen.

When I have concluded the experiments in hand, and have been able to tabulate my results, I shall make known the method.

Laboratory, Upton Manor, Plaistow, E.
July 1, 1879.

PREPARATION OF AMMONIA-FREE DISTILLED WATER.

By J. S. THOMSON.

THE method of water analysis as elaborated by Messrs. Wanklyn and Chapman leaves but little to be desired in the matter of simplicity. The only part of the process which is in any degree tedious is the preparation of the ammonia-free water. That such is the case is, I think, clearly shown by the number of suggestions which have from time to time been made with the object of either doing away with distillation in its preparation altogether or of at least securing a larger proportion of the distillate fit for use in the process of analysis than is usually the case. These various processes, although no doubt very ingenious, have never come into general use, chemists still preferring to prepare the water by the original process, tedious though it be. The process about to be described appears to be free from the objections commonly urged against the earlier ones, and at the same time it has the advantage that the original cost of the necessary apparatus is but small.

The following modification of the distilling apparatus commonly found in laboratories at once provides a simple means of securing a practically unlimited supply of ammonia-free water and at the same time demands little or no attention. The tube which conducts the water vapour from the still or boiler in place of being connected directly with the worm or condenser pipe, is made to enter at the bottom of a large iron drum (say 10 gallons), while the top of the drum is connected with the worm pipe in the ordinary manner. The drum, which is freely exposed to the atmosphere, condenses a portion of the aqueous vapour, which, falling to the bottom, is kept in a state of ebullition by the free steam blowing through it. Now, if this be drawn off by a suitable tap it will be found to be perfectly free from ammonia, while the ordinary distilled water got at the worm end is teeming with ammonia. With apparatus of the dimensions given above, with a plentiful supply of steam, the ammonia-free water can be collected at the rate of about one gallon per hour.

Addiewell Chemical Works,
June 30, 1879.

NATURE OF COHESION AND ITS CHEMICAL SIGNIFICANCE.

By FRIED. MOHR.

AMONGST all the properties of matter cohesion is the most universal in its manifestations and yet the least regarded. We are so accustomed to perceive all things in a certain state of cohesion, that we cannot separate this attribute from the essence of things, whence it happens that it has been treated as the Cinderella of physics and

seems as if it were placed outside the law of the conservation of force. An attempt has been made to explain the phenomenon by means of the attraction of the smallest particles, and to bring this attraction into connection with the universal attraction of the heavenly bodies, which is also merely an assumption. It is plain that cohesion is a true force which can be measured by the sum total of force required to overcome it. We must endeavour to bring this cohesive force in connection with the other accessible and known forces according to the laws of mechanics. The force most readily to be connected with cohesion is heat, because we know that heat can overcome cohesion. We must seek to understand how heat—which we rightly consider as an internal movement in substances—exists in such substances, and for this purpose the theory of waves offers us the needful points of approach.

After summarising the characteristics of undulation, and distinguishing progressive and stationary waves, the author proceeds:—

Cohesion is not due to thermic vibrations, as, on the contrary, heat annuls cohesion. A second force must therefore be present which produces the phenomena of cohesion. Thermic vibration has merely served to explain the mechanism of internal motion.

The existence of a second force or movement present in all bodies, and essentially distinct from heat, is inferred from the heat which appears on chemical combination. We must always remember that force is never created or annihilated, but that in all cases there occurs merely a modification in the form of the movement, but not in sum of the *vis viva*. If, therefore, heat appears in a chemical process, it must have been present in the reacting matter in some other form—as motion which is not heat. For storing up a great force in a small quantity of substance no other form can be found save that of stationary waves, which cannot become progressive; and in solids we assume a vibration much smaller than that of the heat waves as regards its volume, but much greater as regards the number of the waves. For this movement I proposed, in 1868, the name "chemical motion," or chemical wave.

We see everywhere that heat overcomes cohesion, and in so doing disappears as heat. This state has been termed latent heat, though we must remember that it is no longer heat, but a movement of another kind. If mechanical power is produced by heat, such heat likewise becomes latent, and the movement of masses, like the thermo-electric current, may be called latent heat.

A proof that a great sum of *vis viva* exists in bodies, in addition to heat, may be given experimentally by means of the calorimeter.

The author adduces experiments in proof of this proposition, and continues:—

Everywhere cohesion is modified by chemical processes, and inversely in every chemical process a change of cohesion appears.

After considering the action of cements, solders, and of the apposition of smooth clean homogeneous surfaces, Dr. Mohr continues:—

Between cohesion and adhesion there is no difference; cohesion was sometimes said to exist between similar and adhesion between dissimilar bodies. If a thick solution of glue is dried up in a porcelain capsule, pieces of the latter may be broken if the glue is removed forcibly. Here adhesion is stronger. Cohesion is a decidedly chemical property, and can be overcome by chemical means. Solution is the fusion of a solid body in one already melted. In the latter its peculiar molecular motion which determines its chemical attributes has been already overcome or modified by an excess of heat, a part of which has apparently disappeared as such, and has been converted into another form of movement. The excess of heat still suffices to overcome cohesion in another body, and thus again a certain quantity of heat disappears as such. Latent heat is not heat, but a new chemical attri-

bute. The 79 heat-units which a weight-unit of hot water at 79° conveys suffice to overcome the cohesion of one weight-unit of ice at 0°, and to convert it into water at 0°, and as long as the water remains liquid no other free heat is present than that which belongs to it as water. But as soon as the water is made to freeze 2×79 heat-units escape, and there remain two weight-units of ice at 0°. This leads us directly to the connection between hardness and fusibility. The latter is decidedly a chemical attribute, and their connection secures the same rank for hardness. The metals form a scale of hardness almost in accord with their fusibility. The fats form two parallel scales of hardness and fusibility from white wax to oil of almonds.

It has been already shown that cohesion consists essentially in the perflux of the stationary waves of that movement which permanently determines the chemical attributes of bodies, and we distinguish the permanent non-communicable movements from the transient movements of free heat. Every permanent action of heat which reveals itself by a change of cohesion is attended by a permanent change of the chemical nature. [?]

Cohesion and affinity being closely connected we are led to consider more closely the views on affinity. The ordinary doctrine is that affinity exists only between bodies which are chemically different. This is correct if we rank among the manifestations of affinity merely such processes as are attended with explosion, ignition, or the production of a strong heat. But it seems that we might exactly invert this proposition, and, as in the animated world, assume affinity among like bodies, if we suppose that the mobility of matter depends on heat. Thus water unites with water in every proportion without any perceptible physical phenomena. But water with its 88·9 per cent of oxygen has no affinity for liquids poorest in oxygen, such as benzol and petroleum. Ethereal oils dissolve in water almost proportionally to their percentage of oxygen, and phenol is tolerably soluble. Potassic chloride and iodide, ammoniac chloride, &c., dissolve in water, on account of the analogy between chlorine, iodine, and oxygen. Between water and fats there is not even adhesion or moistening. The internal movements are respectively so heterologous that the wave-systems cannot pass into each other; on the other hand, liquid oils adhere to solid fats, as does mercury to gold.

It was observed that certain chemical bodies can be mutually substituted for each other in crystals without a change of form. This fact was named isomorphism. When it subsequently appeared that the atomic volumes of isomorphous bodies was equal, the idea of isomorphism was transferred from the crystal to the atom, it being assumed that the substitution was explained by the equal size and similar shape of the atoms. Thus sulphur, selenium, chromium, and manganese appeared isomorphous on account of the acids RO_3 , and manganese and chlorine on account of the acids R_2O_7 . If isomorphism depends on the identical form of the atoms, chlorine must be isomorphous with sulphur, selenium, and chromium. This is not the case, and the assumption is therefore false. Sulphur has two crystalline forms differing in specific gravity, and consequently in atomic volume, which is atomistically impossible and unthinkable. The acicular sulphur when it passes into the rhombic form by comminution evolves heat, and on burning evolves 41 calories more than does the rhombic variety, so that this heat must have entered into crystallisation. The assumption of the equal shape and size of atoms is therefore untenable.

Our modern theoretical chemistry depends entirely upon the atomic hypothesis, i.e., it is assumed that the elements consist of smallest particles not further divisible, and hence called atoms. On this point we have neither direct observations nor experiments. The assumption is made in order to explain the fact, a thousand-fold confirmed, that the elements always combine with each other in definite ponderable proportions, and that if two

elements form several combinations, their proportions are small multiples, not exceeding the number 7. In fact the atomic theory explains this phenomenon satisfactorily, but nothing more, and great modesty has been shown in endowing these atoms with properties because important mechanical difficulties cropped up on all sides and weakened the indispensable proof for multiple proportions. We have no idea of the form, the colour, the magnitude of these atoms, and we have merely deduced their relative weight, their relative volume, and their specific heat. Our monistic conception of nature leads us to apply to atoms our view as to the cohesion of bodies, since the assumption of indivisibility must find its ultimate basis in cohesion only.

The magnitude of the atomic weights has been deduced from analyses, leaving merely the uncertainty whether a single or a double weight must be admitted, according as we proceed with or without Avogadro. The atomic volume equals the atomic weight divided by the specific gravity, and the atomic heat is the product of the specific heat and the atomic weight.

The calculations founded upon these propositions depend on the assumption that the specific gravity of the atom is the same as that of the mass. The discussions concerning atomic volumes are well known. According to the experience that the atomic weight of a compound may be found by adding up the atomic weights of its constituents, we might imagine that the atomic volume of a compound would be calculable in a similar manner. This, however, is never the case. The specific gravity of the individual constituents is never the same in the compound as in their free condition, but expansion or contraction always ensues. The compound never occupies the same space as its constituents. Carbonic bisulphide occupies 41 per cent more space than its constituents. Its specific gravity, calculated from that of carbon and sulphur, should be 2·152, but it is found to be only 1·272. Consequently atoms have to be credited with a power of expansion or contraction according to circumstances. The calculation of the atomic volume presupposes that the atom of silver possesses the specific gravity of a mass of silver (10·428). We must then also assume that it has the lustre and the opacity of massive silver, not to speak of hardness, because it is indivisible. If we once assume atoms we must grant them not merely indivisibility, but also immutability, and it is then inconceivable how we can see through melted nitrate of silver which contains by weight 63 per cent, and by volume 6 per cent, of metallic silver. The mercury atom must be a minutest granule of mercury, and equally opaque as mercury in bulk. Concerning carbonic acid and ether, we know that if heated in strong glass vessels they disappear, and yield a vapour almost of the same density as the liquids themselves. If we assume this to be the case with mercury, the absolutely transparent and invisible vapour of mercury would have nearly the specific gravity 13. This fact, and, indeed, the changes of all bodies by mere heat, is on the atomic theory incomprehensible. A different arrangement of the atoms cannot possibly deprive them of all their attributes.

The atomic theory meets with its greatest difficulties in the gases. It is necessarily assumed that the atoms of hydrogen, oxygen, &c., are solid, massive, indivisible bodies. The specific gravity of these atoms cannot be determined, since heat and pressure cannot be removed from them. Most of these gases are absolutely invisible, and according to the analogy of mercury and zinc, it would be possible that such atoms also may be of an opaque, metallic nature. But the liquefaction of the permanent gases, lately so-called, yields colourless liquids, which, indeed, like water, are invisible by transmitted light. The atomic theory is compelled to assume that the gases move as distinct particles in an absolute vacuum. But then we cannot understand how light can penetrate. The optical ether is no more a matter of demonstration than the atoms, and, as we deny its materiality, it cannot

be the substratum of a force. The gas theory of König and Clausius is a rather bold attempt to explain the existence of gases; but it cannot maintain its ground even on mechanical principles. It is assumed that the absolutely elastic gas atoms move with equal speed in a rectilinear direction, and rebound from the sides of the vessel. At the same time they vibrate in all directions. It is mechanically inexplicable that a body moving in a certain direction should ever change its direction without some external influence. The individual atoms must share among themselves the function of vibrating in different directions.

If in a cubic metre of empty space there is one cubic millimetre of hydrogen gas, there must be spaces perfectly void. The nature of gases is inexplicable, unless we consider the matter of the gases as continuous and perfectly elastic, whereby the empty spaces are done away with. This applies to all bodies which can assume the gaseous state or can combine with solids according to the law of multiple proportions, *i.e.*, to all the elements without exception. The atomic theory fails to explain the changes produced in bodies by heat alone; the difference of properties in compounds and in their constituents; the liberation of heat which accompanies combination, and its disappearance during fusion and dissociation; it does not explain allotropisms, colours, states of cohesion; in a word, it explains nothing save the uncontested fact of combination in definite proportions.

Atoms which according to theory are solid bodies must have a shape. On this subject nothing has been observed. As the most natural conjecture the sphere has been assumed as the most regular of all forms. But in pursuance of another theory the existence of molecules has been inferred, *i.e.*, combinations of from two to six atoms forming one common body, and it is conjectured that the gases in a free condition consist, not of single atoms, but of molecules each formed of two atoms.

For this view we have no proof save its agreement with a hypothesis itself incapable of demonstration. It cannot be shown how the tendency of the individual atoms towards union can be satisfied by any particular number, since between two individual bodies, *ex hypothesi* absolutely identical, no compensation of different forms of motion can take place. We are unable to say whether a mass of silver or copper is composed of atoms or molecules. Two spheres can touch each other only at one point, and, as we have shown above, no notable cohesion can thus arise. The spheres have also been invested with absolute elasticity, without which the permanence of a gas cannot be thought. If we wish to assume that these spheres flatten each other, and thus offer a larger surface of contact, there can be no reason advanced why the atoms should press so closely together. A body consisting of spheres must necessarily contain empty spaces, into which the finer atoms of gases, such as hydrogen, may penetrate. But not a single fact supports this view (?); and we must, therefore, assume that massive bodies completely fill the space which they occupy.

The origin of a body continuously filling space, as the doctrine of atomic volumes demands, presupposes atoms as bounded by plane surfaces.

The strong cohesion of solids compels us to assume that their mass is continuous, and at the same time elastic, so as to receive the waves of thermic and cohesive movement. If we sum up in brief our results, they may be formulated as follows:—

1. Cohesion depends on pervading stationary waves of a particular form of motion, which at the same time determines the chemical properties of bodies. These waves are not directly transmissible.
2. Heat is another form of movement with both progressive and stationary waves, which can be communicated from one body to another.
3. In the chemical union of two bodies a part of the chemical movement is eliminated as heat, or heat is taken up as chemical movement. In the former

case there is an increase of cohesion, and in the latter a decrease.

4. Heat is everywhere present in the free state, and everywhere it diminishes cohesion.
5. The quantity of *vis viva* which dwells in any substance as its chemical quality exceeds the free heat immensely.
6. Solids have unchangeable nodal planes and stationary semi-waves at their boundaries. Liquids have movable nodal planes.
7. Vapours are gases from which the chemical wave-movements are easily separated as heat. Gases are vapours from which these movements are not readily separable, the difference being merely one of degree.
8. All the phenomena of cohesion can be reduced to vibration.
9. Allotropic forms are distinguished by unequal quantities of chemical movement, and of the accompanying differences in cohesion, as proved by unequal heats of combustion.

The facts of definite proportions and of small multiples must find a new explanation, and if the suspicious word "atom" is to be avoided we may speak of combining weights. The law of Dulong and Petit of the equal atomic weights of the elements, though not thoroughly carried out, does not depend upon accident, and must have a real basis; likewise the fact of similar atomic volumes in bodies chemically similar. The further fact must be recognised that the gases combine in simple proportions, and that their combining weights coincide with their specific gravities. The magnitudes of our atomic weights will also not be influenced by a new theory.

The laws of wave-movement are the same for the revolution of worlds, for the waves of water, the pendulum, the vibrating string, for sound, light, heat, and cohesion.

As the phenomena of cohesion and the chemical properties of bodies therewith coincident depend on the transit of waves, the long-recognised fact becomes intelligible that chemical action ensues only on the immediate contact of the bodies, that the phenomena of affinity do not depend on a specific attraction, and that there can be no question of a static of atoms. Chemical combination is the act of assimilation of different systems of waves, and it is the more intense the more they differ in their qualities.

After chemical combination the two bodies interpenetrate each other completely, and possess one and the same wave-system, but different from the systems of their free condition.

As the colour and hardness of bodies are consequences of their inner wave movements the change in the properties of the new compound are easily explained by the elimination of movement in the shape of heat.

Dissociation is the rupture of a chemical compound by heat which is permanently introduced as chemical movement and becomes latent. One at least of the constituents must be gaseous or be capable of becoming so.

The explanation of definite combining proportions by means of the atomic theory was, further, very difficult, and was not reached without some *sacrificio dell'intelletto*, which, however, was overlooked, as the consequences were not inferred. On attempting to bring cohesion within the law of the conservation of energy all this came to light, in addition to the inextinguishable mechanical difficulties which cling to the idea of the atom. Along with atoms must fall their enumeration, their catenation, their position, the molecules and their "splitting," the structural formulæ, the types, the rings, the unities of affinity, and the whole atomistic *canon* as at present in vogue. Modern chemistry has placed herself outside the law of the conservation of energy; she accepts the heat of combination as a free gift without asking its origin; she explains the different properties of isomeric compounds by a different position of the atoms, though such compounds display different combustion-

heats and from position alone no movement can arise. Lothar Meyer's ingenious syllabus of modern chemical theories begins with the sentence:—"The foundation of all at present prevailing chemical theories is the atomistic hypothesis," and the burning question of the combustion-heats is not noticed in a single syllable; not even the feeling of the need of an explanation of this the most important of chemical processes can be traced. If based upon a false foundation science can make no valid advances, and explanations turn now upon hypotheses rather than upon nature. In due time an explanation of the law of multiple proportions, based upon the theory of undulations, will be found, and instead of a proud *ignoramus* we shall have modest *ignoramus*, or even a hopeful *invenimus*.

The foregoing representation is indeed merely like a blow on water, which stirs up a few ripples, but no persistent ones. It is easier to agitate atoms than to shake the faith in them, and I hear many a voice exclaim, "Disturb not my structure formulæ; are all the time and labour expended in making them look like something real to be wasted? or can you give us a new faith in place of the old?"—*Annalen der Chemie*.

PROCEEDINGS OF SOCIETIES.

PHYSICAL SOCIETY.

June 28th, 1879.

Prof. W. G. ADAMS in the Chair.

NEW members—Mr. J. F. Moulton and Mr. J. J. Eastwick.

Prof. W. G. ADAMS, the President, exhibited his new measuring polariscope. It consists of three principal parts. The lowest section consists of a mirror, a lens, a Nicol's prism, and two other lenses. The upper section consists of lenses and Nicol's prism arranged in the reverse order. Each lens and Nicol's prism is supported separately by screws, and its position can be altered independently of the others. These two parts form a complete polariscope. Besides these there is a middle piece, consisting of two lenses (nearly hemispheres), forming a box to enclose the crystal immersed in oil, their curved surfaces being concentric. The whole middle piece is supported on the tubes of the upper and lower portions, and may be turned about the optical axis of the instrument. The vertical graduated circle carrying the central lenses and crystal may be turned through any angle about its horizontal axis. By means of an arc fastened perpendicularly on the graduated circle, with its centre at the centre of curvature of the central lenses, the crystal may be turned about another horizontal axis at right angles to the former, so that the crystal and the central lenses can be turned about each by three axes which are mutually at right angles. By means of a system of toothed wheels in gear with the rims of the central lenses, the crystal and central lenses may be turned separately about the optical axis of the instrument, so as to bring the planes of the optic axes of a biaxial crystal parallel to the plane of the vertical graduated circle.

Sir JOHN CONROY, Bart., read a paper "*On the Distribution of Heat in the Spectrum*." After referring to Dr. J. W. Draper's supposition that all the rays in the spectrum have the same heating effect, and to his statement that owing to the unequal dispersion of the prism for rays of different refrangibility the method that has been usual for determining the calorific intensity of the various parts of the spectrum is an essentially defective one, the author described a graphical method for eliminating the effect of the unequal dispersion of the prisms, and showed that from MM. Fizeau and Foucault's measurements, and also

from those of Lamansky and Prof. Tyndall, that the maximum intensity is about the middle of the visible spectrum and not at the red end; and, further, that the curves given by various observers as representing the intensity of the heat in different portions of the spectrum are, in reality, the "dispersion curves" for the particular prisms employed.

Captain ABNEY, R.E., called attention to his published paper "*On the Measurement of the so-called Thermo-Spectrum*," wherein he shows that the distribution of heat in the spectrum is a misnomer, and that what was really measured by Lamansky and Tyndall was the energy absorbed by the lampblack and the absorption due to the prisms used. He considered that there was no inherent heat in the spectrum. He found that Dr. Draper had not taken into account the amplitude.

Prof. GUTHRIE said that Captain Abney had expressed what many thought, namely, that heat was radiant energy.

Mr. GRANT then described an investigation which he had made into the induction lines round two parallel coils of wire in the primary coil, an intermittent current of electricity from a Leclanché battery flowed; and in the secondary, a telephone was connected up to detect the induction sounds. With this apparatus he found that with the coils kept parallel to each other, there were lines, or rather a surface of minimum induction, surrounding the primary, and that if the secondary were placed in these lines hardly any induction noise could be detected. A diagram, representing a medial section through the coils, showed the lines to proceed from the wire of the coils in two curves resembling parabolas, one from each cross section of the wire outwards.

Dr. SHETTLÉ then described his experiments proving the lines of force in a bar magnet to run spirally round the bar between the equator and poles, the equator being decentred and oblique across the bar, as shown by diagrams.

Prof. ROWLAND, of Baltimore, made some observations on the new theory of terrestrial magnetism of Profs. Ayrton and Perry. He said the experiments on which the theory was founded had been attributed to Helmholtz, but they were entirely his own, he having gone to Berlin to make them. The new theory had occurred to himself on making these experiments, but he had rejected it because he found that the potential which the earth's surface would require to have would not only cause violent planetary disturbances, but by mutual repulsion drive objects off the earth. He had made also an experiment to see if absolute motion of electricity would cause magnetisation, but failed to get any effect from it. Then he resorted to calculation to find the magnetic effect of relative motion by rotation of a changed sphere of perfect magnetic permeability that is more magnetic than iron. He found that when the sphere was uniformly charged and rotating there would be a magnetic field in its interior; but, instead of the result of Messrs. Ayrton and Perry, that if the earth were charged to a potential of, he believed, 10^8 volts relatively to interplanetary space, the earth's magnetism would be what it is, he found the necessary charge to be $61 + 10^{11}$ volts. In the ordinary atmosphere this potential would produce a spark nine million miles long, and discharge across to the moon. If the moon were electrified to the same degree, the mutual repulsion would overcome the force of gravity between them. He therefore considered terrestrial magnetism to be still a mystery. He had also thought that the aurora borealis might be explained by supposing the upper regions of the earth's atmosphere electrified. The winds carrying the upper strata towards the poles, electricity would condense there. This hypothesis is tenable still.

Prof. AYRTON said that whether or not the new theory of magnetism should be so rejected depended on whether or not Prof. Rowland's calculations, or those of himself and Prof. Perry, were wrong. It had been found by Sir William Thomson, from experiments at Arran, that the earth was electrified with respect to the air, and that

there is a difference of potential of 30 volts between earth and air for each foot of ascent. This gave 1360×10^{11} centimetre-gramme second electrostatic units as the potential of the earth. The new theory required the potential to be 1011×10^{11} , or supposing the earth to be solid iron, or about 14 times more, a wide margin.

Prof. ROWLAND said he had not seen the calculations of Profs. Ayrton and Perry yet; but he believed his results to be correct, as he had checked them in various ways.

Mr. RAILLEY exhibited a modification of Arago's experiment, in which a copper disc is caused to rotate continuously by changing the polarity of four electro-magnets underneath by a revolving commutator.

Mr. CONRAD COOKE exhibited a single voltaic element showing the internal current. This is done by forming the glass vessel containing the element into a helical tube between the poles, and hanging a galvanometer needle in the interior of the helix; the internal current deflects the needle.

NOTICES OF BOOKS.

Supplement to a Handbook of Chemical Manipulation. By C. GREVILLE WILLIAMS, F.R.S. London: Van Voorst. 1879.

TWENTY-TWO years have passed since the author published his excellent handbook. This addition of some 90 pages to the 580 of the original volume is most welcome. We may wish it to have been more complete; we may think some parts of it treated more fully than others; but we are bound to acknowledge that Mr. Greville Williams has, on the whole, combated the difficulties that beset all supplement writers with a very fair measure of success. He has not been hampered by the desire to attain uniformity, but has very properly sacrificed the old notation to the new in the pages now added to his former work, and he freely confesses that he has omitted treating of some large sections of his subject because they could not be discussed adequately within the narrow limits imposed upon him. The following notes may perhaps suffice to show what our author tells us himself, and what he recommends us to seek for in the works he cites.

Under "Furnaces" we have references to the catalogues of Fletcher and Griffin, with an excellent woodcut of the foot-blower devised by the former apparatus designer. Sprengel's specific-gravity apparatus is figured, and its employment described on page 9. Nine pages are given to vapour densities—a favourite subject with the author of this book. The ingenious arrangement for continuous filter-washing figured on page 21 is due to Mr. Greville Williams himself. Another and still more effective piece of apparatus for the same purpose is described and illustrated on page 22. Various pressure filters are noticed on pages 23 to 26. Pressure-tube operations are then described, and afterwards boiling-points, distillations, and melting-points. Under the heading "Volumetric Manipulation," page 38, the use of the Orange No. 3 of Porrier for indicating neutrality in determining acids and alkalis is recommended, examples of and directions for its employment being fully described. This substance is *ammonium dimethylamidoazobenzosulphonat*. For eudiometric improvements reference is made to the memoirs of Russell in the *Chemical Society's Journal*. The next 20 pages of this supplement are occupied with a most instructive and important digest of improvements in organic analysis, especially so far as relates to combustions for carbon and hydrogen determinations; the illustrations accompanying the text are well selected and carefully executed. For many operations in physical manipulation the author refers us to Roscoe's "Spectrum Analysis," and Weinhold's "Introduction to Experimental Physics," but a few pages are devoted to new

forms of galvanic batteries, while an excellent account, with figures, is given of Sprengel's pump.

Altogether the eighty-eight pages of letterpress and the twenty-three illustrations before us are well worth the moderate sum asked for this useful appendix to an invaluable book.

Laboratory Teaching. By C. L. BLOXAM. 4th Edition, London: J. and A. Churchill, 1879.

THE alterations made in this edition chiefly concern the formulæ now for the first time introduced. The work, perhaps, requires some systematic revision, but there can be no doubt that it still merits and still retains its character as a very useful practical handbook for the beginner in qualitative analysis. We should, however, like to see such phrases as the following modified:—"Sulphide of ammonium or hydrosulphate of ammonia, $(\text{NH}_4)_2\text{S}$, is common in a state of solution only" (p. 68). Surely this is awkward. So also it is perplexing to the young student to find another formula, namely, NH_4HS , given for hydrosulphate of ammonia in the index, while the terms ammonium sulphide and ammoniac sulphide are assigned to the previous formula in another part of the index (pp. 232 and 234). The author justifies some of the discrepancies between his formulæ and his names for chemical compounds by the statement "that an endeavour to be absolutely consistent would injure the practical usefulness of so small a book." This is a dangerous argument. A similar reason is assigned for the assertion on page 142 that dextrin is insoluble in cold water, like starch, gum, soap, and the oxalates and nitrates of urea.

On the Estimation of Phosphoric acid by Magnesia for Commercial Purposes. By E. F. TESCHEMACHER and J. DENHAM SMITH. London: Hardwicke and Bogue, 1879.

THIS pamphlet of 32 pages, on the special mode adopted by the authors of carrying out the analysis of commercial phosphates, scarcely calls for remark. The writers confess that they "do not care to claim the replacement of tartaric by citric acid, of sulphate by muriate of magnesia." Their process is, in fact, the ordinary oxalic-acid method, complicated by several elaborations, and involving the *simultaneous* employment of oxalic and citric acid in that stage of the work in which the lime is precipitated. This we consider a mistake. The employment of sulphite of ammonia as an aid in retaining iron in solution has well-known advantages, but is not essential to the complete success of the oxalic acid process. With many of the other directions in this pamphlet we are dissatisfied: such directions are those on pp. 24 and 25, where the ammonia magnesian phosphate has but a single hour given it to form and completely separate from a solution; the employment of no less than 350 grain measures of the magnesian precipitant; and the recommendation of a porcelain crucible for the ignition of the precipitate. We should like to have seen a word about the analysis of those natural phosphates in which more than traces of magnesia is present. In the year 1879 surely the terms *muriate* of magnesia and *muriate* of ammonia (p. 24) are a trifle antiquated!

Two New Isomeric Cyanuric Acids.—J. Herzig.—The α -acid differs from the normal cyanuric acid by crystallising with 1 mol. of water instead of with 2. 100 parts of alcohol dissolve 0.349 of the normal acid and 0.556 of the α compound. The β -acid is much more soluble in alcohol and water than either of the above, and if heated in a glass tube it does not evolve vapours of cyanic acid.—*Berichte*.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 65, May, 1879.

Report by M. Aimé Girard on M. Kuhlmann's (junior) Methods of Conveying Acids.—M. Kuhlmann, in place of carboys, employs floating reservoirs in the form of an ordinary boat, fitted with air-chambers to give them sufficient buoyancy. For sulphuric acid of 60° B. and upwards these are constructed of sheet-iron, and have been in successful use for some years on the canals of the North. For hydrochloric acid he uses cylinders of hardened india-rubber, kept in their form by an external framework of wood. A modification of the structure serves for transport by rail.

Report by M. Troost on M. Gaiffe's Galvanic Deposits of Cobalt.—The metal is deposited from a solution of the double sulphate of cobalt and ammonia, and is superior to nickel at once in hardness, tenacity, and in beauty of colour. It is much less oxidisable than iron, but is very easily dissolved by acids.

On the Production, Constitution, and Properties of Chrome-Steels.—M. Boussingault.—An extensive memoir, taken from the *Annales de Chimie et Physique*, the first portion only appearing in the present issue.

Biedermann's Central-blatt.

No. 5, May, 1879.

Prediction of the Weather.—Prof. Klinkerfuss.—The author gives the following conclusions:—The dew-point read off at sunset indicates the minimum temperature for the night. If the dew-point is below 0°, and the sky clear, frost is to be expected. If it is about 5° below the mean temperature for the day, a cold and dry current of air is approaching. Inversely, a high dew-point indicates that a warm and moist current is approaching, when the air may easily reach its point of saturation, rain then being the result. If, in a day hotter than the average, the dew-point equals or exceeds this temperature, thunderstorms may be expected, on account of the quantity of condensed vapour. If the dew-point rises to 20° C., hail may be expected.

Formation of Nitric Acid in Soils.—Prof. Hünefeld.—The author maintains that the higher oxides of manganese, along with magnesian carbonate in contact with water and due access of air, have the power of inducing the formation of nitrites, and subsequently of nitric acid. Affirmative results have been obtained after the materials had been previously found free from oxides of nitrogen.

Examination of the Shells of Molluscs and Crustaceans.—Prof. F. H. Storer.—The shells of molluscs may be regarded as agriculturally worthless, but those of crustaceans contain notable proportions of phosphoric acid, potash, and nitrogen.

Has the Texture of Superphosphate an Influence upon its Efficiency?—Dr. P. Wagner.—The author examines whether of two superphosphates, of equal percentage of soluble phosphoric acid, the one most finely powdered is in all cases the most effectual? He concludes that, if superphosphate is applied to a moderately moist calcareous soil, the absorption of phosphoric acid takes place so rapidly that after a few hours almost the whole is rendered insoluble without having been previously distributed to any extent through the soil. Hence it is requisite, especially on such soils, that the superphosphate should be in the finest possible state of comminution.

Die Chemische Industrie.

No. 4, April, 1879.

This issue consists almost exclusively of discussions on the new tariff as affecting chemicals.

According to the *Breslauer Zeitung* considerable deposits of sulphur are found in the gypsum formation of Upper Silesia, especially at Paschow and Kokoschütz, near Ratibor.

No. 5, May, 1879.

The Patents-Committee of the German Association for Promoting the Interests of Chemical Industry (why, we beg to ask, must German associations assume such intolerably long names?) has held a session at which an interesting communication from the scientific staff of the Baden Aniline Works was read. The authors point out, not apparently with satisfaction, that neither a novel substance nor a novel application of a substance obtained chemically can be patented in Germany. Concerning the preliminary investigation they doubt the possibility of deciding *à priori* on an invention.

Aluminium Sulphate as a Disinfecting Agent.—A. Tedesco.—The author is of opinion that aluminic sulphate has been recently proposed as a means of disinfection, for which purpose he considers it eminently adapted. He ascribes to it the following action:—The ammoniacal products of decomposition are fixed as ammonium sulphate; the liberated aluminium hydrate carries down all suspended particles, forming with them a solid precipitate. The organic cell, in contact with aluminous compounds, absorbs alumina with great avidity, losing thereby its vegetative power, and putting an end to the process of decomposition. He considers bauxite and wochenite the best materials for the preparation of a sanitary sulphate of alumina. Kaolins are readily attacked by sulphuric acid, but are poor in alumina and comparatively costly.

Moniteur Scientifique, Quesneville.

June, 1879.

A great part of this issue is taken up with a discussion on "septicæmia and germs," which appears to have engaged the attention of the Academy of Medicine on March 11 and 18 and May 6 and 13.

Influence of Changes of Temperature on the Deviation which Inverted Sugar produces on Polarised Light.—Paul Casamajor.—See *CHEMICAL NEWS*, vol. xxxix., pages 212, 234.

Acceleration of Tannin by means of Phosphoric Acid.—E. Ador.—Phosphoric acid prevents the precipitation of albumen by tannin, but it is without action upon gelatin. Hence it enables the tannin to penetrate the hides more rapidly, and renders it possible to use stronger tan-liquors, but there is a loss of weight if the liquors are stirred.

Reply to Prof. F. Selmi's Remarks on the Toxicological Detection of Arsenic.—A. Gautier.—The author contends that his method is not open to objections on the score of the volatilisation of arsenious acid.

Textile Colorist. Vol. i., No. 4, April, 1879.

This issue contains a great quantity of useful matter, but no novelties of general interest.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 5, May 29, 1879.

The statement has been made in M. Figuier's *Année Scientifique* that M. le Vicomte de Vergnette-Lamotte had discovered a photographic method of analysing wines. The only truth in the rumour is that M. de Vergnette-Lamotte has made some microscopic researches on organised bodies present in decomposing wines, and has caused the results to be represented photographically.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 2, 1879.

Certain Metallic Phosphides.—O. Emmerling.—The author heats metals along with phosphorus in glass tubes exhausted of air, sealed, and bedded in iron tubes filled with magnesia, and then gradually heated to dull redness. The tubes are afterwards opened in an atmosphere of carbonic acid. A phosphide of copper CuP was obtained; the magnesium compound was too unstable to admit of analysis. With aluminium, iron, and mercury there was no action. Other compounds obtained were AgP, Cd₂P, Zn₃P₂, and SnP₂.

Behaviour of Meta-nitro-anisol with Ammonia.—H. Salkowski.—Meta-nitro-anisol is not converted by ammonia into meta-nitro-anilin.

Behaviour of Plumbiferous Concentrated Solutions of Potassium Iodide with Hydrogen Sulphide, and on Plumbiferous Crystals of Potassium Iodide.—E. Schering.—Sulphuretted hydrogen does not precipitate all the lead dissolved in solutions of potassium iodide except they are much diluted. The form of plumbiferous crystals is somewhat modified.

Oxidation of Quinine by Potassic Permanganate.—S. Hoogewerff and W. A. van Dorp.—Among other products the authors obtained a nitrogenous tribasic acid, C₈H₃N₆O₆.

Structural Formulæ of the Aromatic Compounds.—E. Wroblevsky.—Not adapted for abstraction.

Constitution of Iso-diphenic Acid and of Fluor-anthen.—R. Fittig and H. Liepmann.—Not suitable for abstraction.

Determination at High Temperatures of the Vapour-Densities of Substances which attack Mercury.—L. Pfaunder.—This paper cannot be intelligibly reproduced without the accompanying engraving.

Action of Nitrosyl-chloride upon Unsaturated Hydrocarbons.—Paul Tönnies.—The author is studying the action of nitrosyl chloride upon amylene, anethol, &c., and considers that this reaction affords a convenient means for the production of primary amines.

Mono-chloro-lactic Acid and Dichlor-propionic Acid from Glyceric Acid.—M. M. Werigo and Melikoff.—The authors have obtained these acids by heating glyceric acid in sealed glass tubes for three days, with a half or the four- or five-fold volume of hydrochloric acid saturated at 0°.

Boiling-points of the Esters and Ether-esters of the Oxy-acids.—L. Schreiner.—The methyl-ether of an oxy-acid ester boils about 20° lower than the latter; the boiling-point of each ethyl-ether is about equally high with that of the corresponding ester.

On Chrome-blacks upon Wool.—M. Reimann.—In dyeing chrome-blacks upon wool the author considers that chrome-alum—now a residual product from the manufacture of alizarin—may be partially substituted for potassic chromates. He recommends also a mixture of chrome-alum, iron-alum, and tartar, followed up with logwood as yielding a black which combines the advantages of the chrome and of the iron-blacks.

On Dibrom-capric Acid.—C. Hell and P. Schoop.—The composition of this acid is C₁₀H₁₈Br₂O₂; it melts at 135°; dissolves sparingly in cold water, and is decomposed by boiling-water.

INSTITUTE OF CHEMISTRY OF GREAT BRITAIN AND IRELAND.

An Examination in Practical Chemistry in connection with the Institute will be held on Monday, 11th August, and four following days. Examiner—Dr. W. R. HODGKINSON. Candidates are requested to communicate with the Secretary, Mr. CHARLES E. GROVES, Somerset House Terrace, London, W.C.

A Chemist (21), who has had practical experience in works, desires Re-engagement. Good references; no objection to go abroad.—For particulars, &c., address, X. T., CHEMICAL NEWS Office, Boy Court, Ludgate Hill, London, E.C.

A Chemist, who has had several years practical experience, is open to an Engagement. During the last six years has been engaged in the Alkali Manufacture.—Address, W. H. R., Fir Cottage, Weston, Runcorn.

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A Young German Chemist (19), who has studied at a Polytechnic School, and is thoroughly conversant with Inorganic and Organic Analysis, is desirous of meeting with an Engagement in England.—Address, L. E., 8, Harpur Street, Theobald's Road, London, W.C.

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London: TRUBNER and CO., Ludgate Hill.

HYDRAULIC FILTER-PRESSES.

IN the CHEMICAL NEWS of 28th March last (p. 128), which, accidentally, only recently came into my hands, I find an article from Mr. J. Marzell, describing and highly recommending a pretended new construction of the Dehne Filter-press, by which an absolute extraction of the cakes deposited in the press is said to be effected in a novel and peculiar manner. This article contains so many errors and incorrect statements, which may lead to grave deceptions, that a rectification appears desirable in the common interest.

As a characteristic mark of the author's want of experience in the practical working of filter-presses in general, and particularly of presses of the especial construction he describes, I may point out that he maintains, that the so-called new arrangement for the absolute washing of the cake, and extraction of its soluble parts, prevents the air from remaining in the cake, and that in the old construction of Dehne, and in the presses of other makers, such air prevented the perfect penetration of the liquid through the cake, and that the absolute extraction obtained in the new press is principally due to the arrangements provided for the expulsion of the air. Anyone who has worked with filter-presses knows that only very few pulps which have to undergo the process of filtration in a filter-press do contain any air, and that in the way such pulps are brought into the press, by means of a Montejus, or a force-pump, no air bubbles can form in the cakes. Such air bubbles, therefore, only exist in the imagination of the author of your article.

Instead of proving the presence of air in the cakes of Dehne's old presses, and presses of other makers, as should have been done in advancing an assertion never heard of before, the existence of air is simply assumed from the necessity, for instance in the porcelain and clay industry, of further preparing the cakes delivered by the press, by clay cutting machines, before they are fit to be worked on the wheel. Up to this time porcelain manufacturers never had the slightest idea that this was done to remove the air contained in the cake, but were rather of the opinion that the subsequent preparation on the cutting and kneading machinery was necessary for obtaining and mixing a perfectly homogeneous material of sufficient plasticity. It is hardly likely that they will replace the cutting machinery by the new washing arrangements of Dehne's press, which, according to Mr. Marzell's explanations, effect the complete removal of the air, unless this new effect is proved to be something more than pure fancy.

These air bubbles—assumed to be present in the cakes of all filter-presses, but which, except in very few instances, exist in reality only in the imagination of Mr. Marzell—play, in the further explanations of the functions of the new press, and in what is represented to be perfect extraction, a contradictory and most fantastical part. Mr. Marzell infers or assumes that the air bubbles become so grouped together as to form a kind of network of small canals, through which, in the old presses, the washing liquid will find its way without washing the more compact parts of the cake.

According to Mr. Marzell's own statement the essential effect of the so-called new washing process of Dehne is obtained by the air being expelled at the commencement of the operation, and its space filled with the liquid. Nobody who knows anything about the laws of hydraulics will be able to understand why the bubbles presumed to be filled with air in the old presses, and grouped together so as to form a network of small canals, will offer an easier passage to the wash water, and, according to Mr. Marzell, will limit the effect of the washing to the surface of the small canals only; whereas in the new Dehne press the air being expelled out of such small canals in the beginning of the lixiviation operation, and the same being filled with liquid, they should now prevent the easier circulation of the liquid, and force it to make its way through the more compact parts of the cake, and so effect a perfect extraction of all soluble parts. According to the law of hydraulics, fluids under pressure take the

way of least resistance, and after these laws the internal working should be quite the contrary to what Mr. Marzell maintains it to be. The air remaining in the canals should prevent the washing liquid from passing them, and cause it to find its way through the more compact parts, so as to effect a more perfect extraction; and in the Dehne press, always following Mr. Marzell's deductions, the small canals, after the expulsion of the air, should facilitate an easy passage of the liquid, and bring the surface of such canals only in contact with the same, rendering the extraction of soluble particles contained in the compact part of the cake impossible.

Such a deduction, which unwittingly proves the contrary of what it is meant to prove, can only be explained by a total, perhaps pardonable, want of knowledge about the laws of Nature. But less pardonable does it appear that the representative of an apparatus does not possess the most shallow knowledge of the functions of the same, as may be demonstrated by the following remarks of Mr. Marzell. In describing the subsequent ways of the washing liquid, in order to explain the superiority of the Dehne press over all others, he says that the water rises at both sides of the cake in the grooves of the press plate, that on its way it expels the air, and that it finally enters the cake from *both sides*; that by this means it has only to make half the way through the resisting cake in comparison to other presses, where, coming from *one side only*, it had to pass the whole cake; or that in Dehne's new press, taking the cake to be 1" thick, it has to make only a way of $\frac{1}{2}$ " against 1" in other presses. In describing the ways taken by the washing water, Mr. Marzell, after having followed it up to the middle of the cake, has only forgotten to tell us where it now remains. The water from both sides meets in the middle line of the cake under equal pressure and having no outlet, no circulation (which alone can effect the extraction of the soluble particles) can take place—always granted, of course, the process to be such as Mr. Marzell describes it.

The designer, however, of Dehne's press has had more foresight than Mr. Marzell, who, in fact, has not understood either its construction or its work. In the described Dehne press the wash water, exactly the same as in all other filter-presses, is introduced by a corner channel or tubular passage in the filtering space formed between the grooves of the press plate and the perforated metal plate to one side of the cake only; it has to penetrate the whole cake and is led away from the filtering space on the opposite side by a similar tubular passage [see Specification of Dehne's press, No. 1957, Henry Edw. Newton, May 15th, 1878, page 3, line 45, &c. :—"The object of this construction is to introduce the washing fluid by the tubular passage (in the lower projecting pieces) into the filtering space on one side of the cake, and to cause the washing fluid to penetrate the cake in order to fill up the entire filtering chamber, and to make the washing fluid to escape through the filtering space on the other side of the cake into the tubular passage in the upper projecting pieces, &c."]

There is no difference whatever between Dehne's new press and the presses of the other makers, in respect to the way in which the liquid is intended to penetrate the cake and to effect the extraction.

These explanations of Mr. Marzell's, which can hardly be taken for anything else but a mixture of misunderstandings regarding the construction of the press represented by him and its functions, and perhaps of delusive information, Mr. Marzell pleases to call the theory of the press, and he passes on to give information about the result of the filtration and extraction, by quoting results obtained in a sugar manufactory in Bohemia or Moravia, information which no doubt issues from a similarly reliable source as his knowledge about the functions of the apparatus. These results, however, may be judged by the simple fact that the liquor is called pretty *concentrated*, notwithstanding its dilution by the addition of 176 litres (39 gallons) of wash water to the filtrate of 157 kilos. (315 lbs.) of cake.

After having shown the imaginary advantage the Dehne press does *not* possess against older apparatus, and how

far its working does *not* differ from that of the older apparatus, I will now in a few words dwell on the differences that really do exist, and the importance of what are supposed to be improvements.

The Dehne press has two additional so-called tubular passages arranged in the corner of the frames for the circulation of the washing liquid and the expulsion of air, instead of the one of older presses, serving in similar manner for the same purpose, viz.—

1. A second corner channel in the lower part of the frames, which communicates with, say, for instance, the 2nd, 4th, 6th, &c., frame, and with the one side of the cakes, whilst the corner channel in the upper part of, say, the 1st, 3rd, 5th, &c., frames used in the Dehne, as well as in all other presses, communicates with the opposite sides of the cakes.
2. On the same frames, containing in their lower part the above-mentioned second corner channel, he has a third channel for the escape of air arranged on the upper part of the frames, which communicates by branch passages with the same side of the cake as the said second channel.

The purpose of these two additional channels is imagined to be as follows:—Whereas in the old presses the wash liquid was introduced by the channel in the upper part of the frames from above to one side of the press cake and had to penetrate the same, and find its exit through an open cock at the other side; the fluid in the Dehne press is introduced by the corner channel in the lower part of the frame; in rising in the space provided between the grooves of the press plate and the perforated plate, it expels the air contained therein by the third channel provided at the upper part of the same frame, and after the escape of all air, the cock of the air channel being turned off, it is imagined to be driven by the applied pressure diametrically through the cake.

I shall further on dwell more fully on the probability of a uniform *penetration* of the liquid through all parts of the cake, which is the necessary condition for a quick and absolute extraction with the least quantity of water, but anyhow the water will arrive at the other side of the cake, and from there find its exit by the corner channel at the upper part of the frame.

The old presses, therefore, performed the extraction of the soluble parts of the cake exactly in the same manner as these presses of Dehne. It was proved, however, by careful examination of the wash liquid, and the extracted cakes (see *Zeitschrift des Vereins der Rübensucker-Industrie*, &c., 1864, p. 642; *Fahresbericht über die Fortschritte der Rübensucker-Fabrication*, &c., by Dr. Stammer, 4, p. 144; *Walkhoff, Der practische Rübensucker-Fabricant*, IV. Edition, vol. ii., p. 42; Dr. Stammer's *Lehrbuch der Zucker Fabrication*, pp. 444-471), that the cake is extracted on the surface only, the particles contained in the liquid, and apparently taken up from the cake, are in reality washed from the filter-cloths, the interior surfaces of the press plates, of the frames and corner channels, and but very little was taken up from the surface of the cake. The apparent extraction was shown therefore to be of no practical advantage, as the quantity of soluble matter taken up from the surface of the cakes was of too little importance to compensate for the disadvantage of the greater dilution of the filtrate, and the washing of the cloths, and the interior parts of the machine is no gain, as at the next charge they will again be soaked by filtered liquor.

The failure of the process of extraction in all these presses is explained by the fact that the cake being held in a flabby and elastic cloth, the wash liquid finds easier passages through which to arrive to the exit side of the cake, than by going diametrically through the same. It will avoid, therefore, the great resistance of the cake itself, and naturally choose the easier penetration round its corners and edges, and thereby only wash the surface of the cake, and (very unnecessarily) also the filter cloths, press plates, frames, &c. The manner of forming the cake within the press, and supporting it, being exactly the same in Dehne's press as in all the old ones, there is no reason why in Dehne's press the liquid should choose

to go through the compact cake, avoiding the easier passage round the corner and edges.

But another question arises; that is, whether the new so-called "improvements" of Dehne might not perhaps be capable of effecting a real extraction by other means than by the uniform penetration of the liquid through the cake. And this, in fact, appears to be not at all improbable. The liquid passing round the corner and edges soon surrounds the whole cake, and by continuous supply and outflow, an uninterrupted circulation is kept up during the whole time of the operation. In this way an interchange of the soluble filtrate in the cake, and a more or less perfect extraction, is gradually effected by means of a process of diffusion. Such process of diffusion being, however, as is generally known, an exceedingly slow chemical process, it is natural that the time for obtaining a really absolute extraction is long and varies very much, proportionately to the nature and composition, thickness and density, of the cake. It is at any rate much longer than the extraction will take in an apparatus which, by its peculiar construction, guarantees with absolute certainty a uniform penetration of the liquid through the cake. Also the wash liquid obtained by diffusion, as explained, is necessarily very much diluted. These circumstances greatly reduce the practical value of the so-called improved arrangements of Dehne, and render them illusory for most purposes.

Besides these imperfections of the improved Dehne press, it possesses, together with most other presses, the great disadvantage arising from the use of filter cloths. The first cost of such cloths, together with the cost of preparing the filters and the changes, which are generally far more frequent than Mr. Marzell asserts them to be, amounts as a rule, in a German sugar manufactory in one "campagne" of about 120 working days, to the cost of a new press. Many pulps are, if capable at all, at least very difficult of filtration in a filter-press with cloths. The use of cloths, and also the light construction of the presses now generally adopted, only permits very slight pressure; indeed, hardly more than 30 pounds on the square inch. The cakes, in consequence, possess but little consistency, and contain far more filtrate than if they had been thoroughly pressed out, and as for the washing operation, they naturally also require a larger quantity of wash liquid. A further very serious disadvantage of filterpresses with cloths—a disadvantage more specially felt in chemical works—is that the cloths get damaged if employed for the filtration of pulps containing acids or alkalies; so much so, in fact, that they become too expensive to permit certain operations and processes to be carried out by means of such filter-presses.

The Press invented by me and patented in Great Britain a considerable time before Dehne filed his patent, does away with these drawbacks (see specification No. 850, 2nd March, 1877). The filtration and the extraction is effected through solid indestructible filter-media and in a far more perfect way than in Dehne's press. Dehne's so-called improved arrangement for the absolute extraction of the soluble parts of the cake, in so far as it consists in the application of two additional corner channels, constitutes in fact an infringement upon my patent for a press without cloths, which provides in its claims for similar arrangements for the introduction and exit of the liquid, and all purchasers and users of the Dehne press will therefore make themselves liable to legal prosecution.

It is too long to give here a detailed description of this new filter-press without cloths. It has, however, lately been introduced into Great Britain, and

MAY BE SEEN at the Warehouse of MESSRS. B. H. REMMERS and CO., in GLASGOW, General Agents for the Licensees and Manufacturers, the Sangerhausen Actien Maschinen Fabrik and Eisengiesserei (late Hornung and Rabe), in Sangerhausen, province of Saxony, Prussia.

(Signed) Dr. DREVERMANN.

BERLIN, June, 1879.

THE CHEMICAL NEWS.

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NORWEGIUM, A NEWLY-DISCOVERED METAL.

THIS metal has been detected and isolated by Dr. Tellef Dahll in a sample of copper-nickel from Kragerø, in Skjærgaarden. The colour of the pure metal is white, with a slight brownish cast. When polished it has a perfectly metallic lustre, but after a time it becomes covered with a thin film of oxide. It can be flattened out in an agate mortar, and in hardness it resembles copper. The melting-point is 350°C ., and the specific gravity 9.441. Its equivalent appears to be 145.9. Only one oxide, NgO , has been obtained. With sulphuretted hydrogen it gives a brown sulphide, even in strongly acid hydrochloric solutions, which re-dissolves in ammonium sulphide. With a slight addition of potassium ferrocyanide it gives a brown, but with larger proportions a green precipitate. The sulphuric solution is turned brown on the addition of zinc, and the metal is deposited in a pulverulent state. The solutions of this metal are blue, but become greenish on dilution.

We are indebted for this notice to the courtesy of Dr. Dahll, who is continuing the study of the new substance.

ON SOME NEW FORM OF APPARATUS.

(SECOND PAPER.)

By M. BENJAMIN, Ph.B.

A FEW weeks ago, under the above title, I described a water-bath* with constant level attachments (CHEMICAL NEWS, vol. xxxix., p. 160). That such an apparatus was desirable is evident from several expressions of opinion that have been received by me since its description. In one case I was informed that the bath was allowed to run for forty-eight hours without replenishing the reservoir, thus demonstrating its utility.

The apparatus to which attention is now requested is the out-growth of several suggestions made in reference to the application of the constant level attachment. It is a drying-bath, made of copper, with double walls; having a tubulature in one corner, into which a thermometer may be inserted. By this means an even temperature can be maintained; or, in case the bath is filled with oil, that the heat may be raised to any desired point. On the top is an opening fitted with a series of four concentric rings, which permits the apparatus being used as a water-bath, and, as in the engraving, a porcelain capsule may be heated, while the lower portion is employed to dry filters or to make determinations of moisture. The constant level attachment is shown to the left of the figure, and is manipulated in the same way as the water-bath. Within the oven are two tin shelves, each of which is pierced with four holes to support funnels.

* In the CHEMICAL NEWS, vol. xxxix., p. 193, a note by Dr. Korn is inserted. As he takes exception to my descriptions, permit me to call his attention to the following facts. While it is quite true that in the *Zeitschrift* for 1872 the constant level attachment is described, it is also quite true that as far back as 1864, in the catalogue of Noellners, the same apparatus is figured (p. 78, fig. 873), but in form it resembles the one described by me about as much as one carbonic acid apparatus resembles another. With regard to the burette clamp, I regret my inability to find its description in either Noellners (1864), Leybold (1872), Griffin (1877), or in any American catalogue. Also I beg that Dr. Korn will observe that the clamp works by a spring and not by a screw; this property distinguishes it from all previous forms.—M. B.

The dimensions of the apparatus are as follows:—Height, $8\frac{1}{2}$ inches; depth, 6 inches, and width 6 inches. In the door I have had placed a sheet of transparent mica; this, I trust, will prove desirable, for, by simply looking in, one may see the conditions of the vessels, without going to the necessity of opening the door.



As both the drying oven and water-bath are indispensable to every analyst, there is reason to believe that some little expense can be saved by combining these two utensils in one, and thus a benefit conferred on the world of chemists.

New York, June 5, 1879.

THE SOLUBILITY OF STANNIC OXIDE IN HYDROCHLORIC ACID.

By A. E. ARNOLD, Assoc. Inst. Chem.

THERE appears to be a considerable amount of uncertainty concerning the solubility of natural dioxide of tin or that artificially prepared, in hydrochloric acid.

W. A. Miller,* W. Phillips,† Greg and Lettsom,‡ R Fresenius,§ and "Watts's Chem. Dict." assert the oxide to be insoluble in acids, while J. D. Dana|| and L. Bombicci,¶ in their two recent books on mineralogy, observe that it is slightly soluble.

If cassiterite in fine powder, or strongly-ignited stannic hydrate, is digested with strong hydrochloric acid at 100°C ., a very perceptible quantity of tin enters into solution as per-salt. The action takes place more energetically above 100°C ., when the dioxide is heated in a current of HCl gas. In the complete analysis of cassiterites, therefore, this fact should be borne in mind. As an example, there was found in a specimen of the mineral—

Oxide of tin, insoluble..	..	38.49 per cent.
" " soluble ..	..	0.06 "

In a few rare minerals oxide of tin is soluble in considerable quantities in dilute hydrochloric acid, evidently owing to the presence of a stannate of some basic oxide.

The French Association for the Advancement of Science.—The eighth session of this Association will be held at Montpellier, commencing on August 28, the President being M. Bardoux, late Minister of Public Instruction.

* "Inorganic Chemistry," sixth edition, 1878.

† 1852.

‡ "Mineralogy," 1858.

§ Sixth edition "Quant. Anal."

|| "Mineralogy," 8th edition, 1879.

¶ "Course of Mineralogy," 1875-77.

CERTAIN

CHARACTERISTIC COLOUR-REACTIONS
PRODUCED BY THE ACTION OF
AROMATIC HYDROCARBONS AND THE VEGETO-
ALKALOIDS ON FUSED ANTIMONY AND
BISMUTH TRICHLORIDES.

By WATSON SMITH, F.C.S., F.I.C.

ANTIMONY TRICHLORIDE.

Naphthalene.—These reactions are obtained as follows:—About 1½ to 2 grms. of pure crystallised antimony trichloride are fused in a small porcelain crucible, and then further heated somewhat more. Then a small particle of the substance is let fall upon the inner side of the crucible, which is afterwards inclined so that the fluid shall just come in contact with the small particle and envelop it. It melts, more or less dissolved by the heated chloride, to which it communicates in certain cases a certain characteristic colouration, which, on setting the crucible upright again, lengthens itself from a mere spot to a long coloured stripe, more easily observed.

On treating naphthalene in this manner, which is not absolutely pure, a more or less rose-coloured tint is produced, often (though the naphthalene seem quite pure and snow-white) a splendid carmine or crimson tint. If, however, the hydrocarbon be absolutely chemically pure no tint appears. This is, then, a most excellent test for the purity of samples of naphthalene. The subsequent formation on cooling of the beautiful rhombic tables of the addition-compound of antimony trichloride and naphthalene, when the proportions have reached something near 3 : 2, is characteristic of naphthalene.

The reactions with other hydrocarbons were now tried:—

Anthracene.—On adding a mere trace of pure anthracene to a little (1½ to 2 grms.) of the fused trichloride at the bottom of a small porcelain crucible, just as with the naphthalene experiment, a clear yellowish green colouration is produced, not destroyed with excess of hydrocarbon. On cooling, colourless needles may be observed, in all probability of an addition compound of antimony trichloride and anthracene.

Phenanthrene.—This hydrocarbon does not dissolve in the fused trichloride nearly so easily as anthracene does, and on solution gives a faint greenish yellow tint. Now as it is a difficult matter to obtain phenanthrene absolutely free from anthracene, it seems not improbable that the faint tint observed may be due to the presence of traces of anthracene. To endeavour to throw light on this point, I obtained three different samples of phenanthrene, the first brown and impure; the second, almost white, and considered nearly pure; the third, beautifully white and crystalline, and considered absolutely pure. These three samples were subjected to the test in question, when No. 1 gave a strong greenish yellow tint; No. 2, a tint greener than No. 1 but fainter; and No. 3 a faint greenish tint. From this gradation I should be disposed to argue that in all probability anthracene may be distinguished from phenanthrene, if both be absolutely chemically pure, by anthracene giving with the antimony trichloride test a greenish yellow colour, and phenanthrene either giving no tint at all or at most a very faint greenish one.

Diphenyl.—Gives no colouration whatever.

Dinaphthyls.—These hydrocarbons give no colouration.

Phenyl-naphthalene (?)*.—Gives not the slightest tint. (Compare phenanthrene which melts at 100°; phenyl-naphthalene melts at 101° to 102°.)

Stilbene.—For this reaction to proceed well, the fused antimony trichloride must be only so hot that the small crucible containing it can just be supported and held in the fingers. If the smallest speck of the substance now be

allowed to fall on the side of the crucible, and the latter be inclined so as to bring the fused mass in contact with the small particle, this is at once turned a deep red or orange-red, and a streak is observed on righting the crucible, extending down its side, of a brick-red or orange-red colour. This tint easily disappears on stronger heating, even if a tolerable excess of hydrocarbon has been used. It thus differs from naphthalene, whose tint is also quite a different one (rose-coloured or crimson). Thus a trace of rose colour observed in applying the test to a sample of stilbene would indicate the presence of naphthalene, especially if on further heating a portion of the colour disappeared, leaving a faint rose tint.

Triphenylmethane.—Gives no colouration. With excess a greenish tint.

Chrysene.—Professor Graebe, of Geneva, kindly furnished me with samples of the hydrocarbons chrysene and pyrene almost quite pure. They were both re-crystallised, chrysene from benzene, and the pyrene from light petroleum spirit. A small speck of the chrysene is let fall on the side of the crucible, and the fluid is then caused to come in contact with it by inclining the crucible slightly. A golden-yellow spot is produced, the chrysene instantly dissolving, and on righting the crucible a yellow streak is observed. Addition of a further and larger quantity of hydrocarbon causes a golden-yellow tint to appear in the trichloride.

Pyrene.—On treating this exactly like the chrysene, it was observed to dissolve with difficulty, giving a faint green or greenish tint. The trichloride requires to be considerably heated to dissolve the hydrocarbon, more so than with the chrysene.

BISMUTH TRICHLORIDE.

Naphthalene.—Gives under exactly the same circumstances as with the antimony trichloride a brick-red colouration, with a shade of crimson, if not absolutely pure; with chemically pure naphthalene, no tint. On cooling first and immediately before solidifying, fine transparent yellow lanceolate needles separate, most probably consisting of an addition compound of bismuth trichloride and naphthalene, analogous to the antimony compound.

Anthracene.—On dropping the smallest trace of this hydrocarbon into the well-fused chloride it at once becomes a purplish black speck, and on further addition a purplish black colouration is produced.

Phenanthrene.—Gives a brown or greenish brown colour, becoming dark on further addition. It is very probable the dark shade supervening on further addition is due to the presence of traces of anthracene in the phenanthrene. The difference from the anthracene reaction is very great.

Diphenyl.—No colouration whatever.

Dinaphthyls.—With iso-dinaphthyl a very faint brownish tint; with the other two dinaphthyls no change.

I have not continued the experiments with bismuth trichloride, as the difficulty of getting it to fuse readily to a clear liquid makes it far less convenient to employ than antimony trichloride. However, the difference in the anthracene and phenanthrene reactions above noted makes it a valuable means of distinguishing between these two hydrocarbons, in conjunction with the antimony chloride test. Table I. shows more briefly the reactions with the foregoing hydrocarbons.

Reactions with Antimony Trichloride and the Alkaloids.

{ **Conine.**—Gives no definite colour reaction.

{ **Nicotine.**—Also no definite reaction.

{ **Morphine.**—In small quantities no change; in larger quantities, faint greenish tint.

{ **Codeine.**—The same as with morphine.

{ **Narcotine.**—The fused trichloride being hot enough to fume slightly. A minute crystal fragment let fall in the clear fused chloride turns at once dark olive-green, seeming black. With further addition the fused mass assumes an olive-green or greenish brown tint.

* This is the hydrocarbon I have recently obtained by passing C_6H_6 and C_6H_5Br through a red-hot tube. As I have not yet made vapour-density determination or combustion I append the query (?).

TABLE I.

<i>Diphenyl</i> —0	<i>Naphthalene</i> —Not quite pure—splendid crimson tint, or rose colour. Absolutely pure, no tint.	<i>Anthracene</i> —Clear greenish yellow, or yellowish green. On cooling, colourless needles formed, of an addition compound. With BiCl_3 —Purplish black	<i>Chrysene</i> —Golden yellow.
<i>Triphenyl-methane</i> —0	<i>Phenyl-naphthalene</i> (?)—0	<i>Phenanthrene</i> —Faint greenish tint if quite pure. With BiCl_3 —Brown or greenish brown.	<i>Pyrene</i> —Green or greenish tint. Hydrocarbon dissolved with difficulty.
<i>Stilbene</i> —Fine orange red, easily vanishing on stronger heating.	<i>Dinaphthyls</i> —0		

TABLE II.

Volatile Alkaloids.	Opium Alkaloids.	"Strychnos" Alkaloids.	Cinchona Alkaloids.	Unclassified.
<i>Conine</i> —0	<i>Morphine</i> —0	<i>Strychnine</i> —0	<i>Quinine</i> —0	<i>Veratrine</i> —(1) and (2)
<i>Nicotine</i> —0	[<i>Apomorphine</i> —0]	<i>Brucine</i> —(1) and (2)	<i>Cinchonine</i> —0	Brick-red.
	<i>Codeine</i> —0	Fine purple-red.		<i>Atropine</i> —0
	<i>Thebaine</i> —(1) red, (2)—0			<i>Aconitine</i> —Bronze-brown.
	<i>Papaverine</i> —0			<i>Santonine</i> —(1) 0, (2) greenish.
	<i>Narcotine</i> —(2) dark olive-green.			With more than trace, (1) 0, (2) dark bluish olive-green.
	<i>Narceine</i> —(1) 0, with only trace.			
	Yellow with little more			

Here (1) means antimony trichloride fused and further heated to a small extent. (2) means antimony trichloride fused and further heated considerably. 0 means no tint produced.

TABLE III.

On Strongly Heating the SbCl_3 close to Incipient Ebullition.

Thebaine.	Veratrine.	Brucine.
Gives no tint, or one disappearing again like a flash as soon as formed. Generally no tint at all is perceived, the vanishing is so instantaneous. Heated more strongly, an olive-greenish brown tint is produced.	Gives a flash of red colour more slowly disappearing than is the case with thebaine. If more alkaloid be added the tint does not entirely disappear, but leaves a reddish hue, turning to a brown if still further heated.	With a trace only, a fine red stain, only slowly disappearing, and if more used, refusing to disappear, and on further continuance of the high temperature passing into a purple-brown tint.

Strychnine.—Not the slightest colouration, even with considerable additions of the alkaloid. The fused trichloride remains clear and colourless.

Brucine.—The smallest possible trace or particle of brucine let fall into the fused chloride produces a beautiful red or purple-red colour. This small though fine red stain shows exceedingly well in the surrounding clear, colourless, fused trichloride, with the pure white background formed by the porcelain crucible in which the test is made. This is a most delicate and characteristic reaction for brucine. The tint is produced distinctly, even if the trichloride be heated almost to boiling. At the point of incipient ebullition the tint turns to a red brown with a purplish hue.

Quinine.—No colour change whatever.

Cinchonine.—No change.

Veratrine.—A red tint, which may be more exactly defined a "brick-red" tint. The tint disappears, if a minute quantity has been added, on heating more strongly, but if not so small a quantity it only disappears partially, leaving a reddish tint in the fluid.

Atropine.—No colour whatever.

Aconitine.—A brown tint, closely approaching a bronze-brown.

Caffeine.—Gives no tint at all.

Narceine.—A trace gives no tint whatever. A larger quantity let fall into the clear fused mass produces a sulphur-yellow spot, approaching greenish yellow. Disappears on shaking or stirring. Excess of alkaloid colours the mass sulphur-yellow, with faint shade of green.

Apomorphia.—Small quantity, no tint. Larger quantity and strongly heating (though not to incipient ebullition), greenish brown tint.

Santonine.—A trace gives no colouration till heated to incipient ebullition of the trichloride, when a greenish tint is produced. A rather larger quantity also at first scarcely produces any change till more strongly heated as just mentioned, when a dark, bluish, olive-green tint is assumed. This reaction is a very delicate one.

Papaverine.—No tint produced.

Thebaine.—With the least trace let fall on side of the crucible, and the same inclined so as to cause contact of the fused trichloride—a blood-red spot, extending to and producing a blood-like streak on righting the crucible; but the temperature of the fusion must be low enough to allow of the crucible just being supported by and held in the fingers, else the stain (or colour) disappears again instantly. On heating more strongly the stain instantly disappears as soon as formed, leaving a perfectly colourless fluid. With a moderate temperature and larger quantity the fluid is tinted red or blood-red: stronger heating causes (even with excess) almost entire disappearance of the colour. Heating to incipient ebullition causes a new tint to appear, viz., an olive greenish brown, or olive-brown, if the term may be used.

Table II. exhibits more plainly and succinctly these alkaloid reactions.

Brucine, Thebaine, and Veratrine.—In order to test the delicacy of the test as applied to the detection of brucine, a mixture was made of strychnine, atropine, cinchonine,

quinine, and aconitine, and to this a small trace of brucine was added, and the whole was well mixed up, so as to give a uniform mass. A little of the mixture, about as much as would lie on a pin point, was now dropped into the fused antimony trichloride, when a small but distinct and beautiful red spot or stain was immediately produced. This proves the method to be of great value in detecting the presence of brucine in that of many other alkaloids, the reaction being in no way prejudiced by their presence. Of course veratrine could not be distinguished by this test in presence of brucine, but brucine could easily be distinguished in presence of veratrine on account of the difference of the reds, their different intensities, and the extreme delicacy of the brucine reaction. The test is made just as with the aromatic hydrocarbons, only more minute quantities of the alkaloids may be taken.

The value of the reactions lies in the fact that those alkaloids giving no reaction leave the antimony trichloride absolutely unchanged in appearance,—i.e., transparent and clear—whereas those giving the reactions immediately and sharply strike a colour, a clear and definite test being obtained. The red tints produced by these bodies are modified as follows, by using a high temperature, and this method may be used for distinguishing them. (See Table III.)

Here, again, these tints are best observed by letting the particle fall on the crucible side, and gradually bringing the fused SbCl_3 in contact.

I am continuing the investigation of these reactions, and hope soon to furnish further results.

Zürich, University Laboratory, July 6, 1879.

DETERMINATION OF NITROGEN IN THE ANALYSIS OF AGRICULTURAL PRODUCTS.*

By S. W. JOHNSON and E. H. JENKINS.

IN 1872 one of us found† that the mixture of caustic soda and caustic lime known as "soda-lime," and used in chemical analysis to determine nitrogen quantitatively, might be replaced by a much more easily prepared mixture of equal volumes of dry sodium carbonate and slaked lime. There is some difficulty in obtaining sodium carbonate suitable for this use. The super-carbonate, which can be readily dried, often contains nitrogen, and the crystals of sal-soda that, after washing, are free from nitrogen, cannot be quickly dried to a fine powder. After various trials the following process of preparing an effective soda-lime was devised:—

Equal weights of sal-soda in clean (washed) large crystals, and of good white and promptly-slaking quicklime, are separately so far pulverised as to pass holes of $\frac{1}{4}$ inch, then well mixed together, placed in an iron pot, which should not be more than half-filled, and gently heated, at first without stirring. The lime soon begins to combine with the crystal water of a sodium carbonate, the whole mass heats strongly, swells up, and in a short time yields a fine powder, which may then be stirred to effect intimate mixture and to dry off the excess of water, so far that the mass is not perceptibly moist and yet short of the point at which it rises in dust on handling. When cold it is secured in well-closed bottles or fruit-jars, and is ready for use.

During the past year a large number of comparative analyses have been made on bone dust, dried blood, fish-scrap, guano, maize-meal, zein or maize-fibrin, and egg-albumin, using soda-lime obtained as above described, and soda-lime either prepared by ourselves according to the directions of Varrentrap and Will, obtained by purchase (Merck's) or kindly supplied by W. M. Harbshaw, Esq.,

Chemist to the New York State Agricultural Society. These analyses have proved that the new soda-lime gives perfectly satisfactory results in all cases when soda-lime can be employed, or, at least, results perfectly agreeing with those obtained by the help of the soda-lime as usually prepared. It is not needful to adduce analyses here in support of this statement, since its truth will appear from the soda-lime combustions to be shortly given, all of which were made with this new mixture.

This soda-lime is to be recommended for the reasons that the materials for making it, sodium monocarbonate and quicklime, are everywhere procurable in a state of purity, the preparation of many pounds of the mixture may be accomplished in an hour or two with little trouble, and the resulting soda-lime is extremely convenient to use, not absorbing moisture in the mixing, and never swelling in the tube to obstruct it on application of heat.

The controversy that has been so actively prosecuted of late years as to the applicability of the soda-lime method of determining nitrogen to the analysis of albuminoid matters, has led us to review the entire subject as far as possible. It is well known to chemists that a number of experimenters—viz., Nowack, Seegen, Nencke, Liebermann, Voelcker, and Musso—have failed to obtain with the soda-lime method as high results on flesh, milk, and similar substances, as by the use of the so-called absolute method, in which the organic body is burned with copper oxide and its nitrogen directly measured in the state of gas. Even Ritthausen, who has stoutly maintained the correctness of the soda-lime method, having employed it in his elaborate researches on the vegetable albuminoids, has very recently admitted that it fails to give all the nitrogen of some of this class of bodies, and has fallen back on the absolute method as the only one to be depended upon.

Since it is a matter of high importance in agricultural and physiological work to use the most exact and especially the most trustworthy methods, we have endeavoured to investigate the correctness of both modes of analysis, to study the sources of error to which they are severally subject, and to learn what is essential to bring out the greatest accuracy they are susceptible of.

We have been led to the conclusion that with the substances above named, both methods when properly worked give nearly accordant results. Dried blood, dried white of egg, and maize-fibrin (crude, obtained by Ritthausen's methods) containing 12 or more per cent of nitrogen, have yielded us from one to two-tenths of a per cent less of nitrogen by the soda-lime combustion than by the absolute method. Our observations satisfy us, moreover, that this discrepancy is no more due to any fault of the soda-lime process than to the errors of the absolute method; errors caused, probably, by the impossibility of removing the last traces of common air from the mixture, either by long-continued transmission of pure carbonic gas or by exhaustion with the Sprengel mercury pump, or by both conjointly.

The following results of analyses by both methods are offered to sustain our assertions. We have not selected these analyses to establish the point, but give all the results we have obtained since learning the best mode of conducting the analytical processes, and they fairly represent what the two methods can accomplish when applied with suitable precautions. (See next column.)

These analyses illustrate a fact which is general in our experience—viz., that the agreement of several determinations made upon one substance is usually closer by the soda-lime than by the absolute method. This fact goes far to show that the soda-lime process is, to say the least, equal in accuracy with the absolute determination.

The full details of the mode we follow in making an analysis by the absolute method cannot be given here. An outline of the process is as follows:—

The substance, mixed with freshly ignited copper oxide, is burned in a long glass tube which has been exhausted

* From the Report of the Connecticut Agricultural Experiment Station for 1878.

† *Amer. Chemist*, vol. iii, p. 261.

	By the Absolute Method. Per cent.	By Soda lime. Per cent.	Average difference. Per cent.
Egg Albumin.. ..	{ 12'54 .. 12'51 .. 12'56 .. 12'44 .. 12'59 ..	{ 12'34 .. 12'36 .. 12'38	{ 0'17
Maize Fibrin (crude) ..	{ 13'73	{ 13'60 .. 11'78	{ .. 0'13
Dried Blood	{ 11'90 .. 11'91	{ 11'72 .. 11'88	{ .. 0'12
"	{ 7'54	{ 7'58 .. 9'12	{ .. 0'04
Fish Scrap	{ 9'21	{ 9'02 .. 8'64	{ .. 0'14
"	{ 8'79	{ 8'71 .. 8'71	{ .. 0'11
Peruvian Guano	{ 8'27 .. 8'63 .. 8'66 8'21 9'68 .. 7'56 ..	{ 8'21 .. 8'49 .. 8'51 .. 8'13 .. 8'04 .. 9'63 .. 9'67 .. 7'56 ..	{ 0'06 0'15 .. 0'12 0'03 .. 0'00

by means of a mercury pump; the products of combustion after passing over ignited metallic copper, and finally over ignited copper oxide,* are collected in a graduated receiver, containing fifty per cent solution of caustic potash, and surrounded by a water-jacket. Complete combustion is ensured by heating, in due course, several grms. of potassium chlorate at the rear of the tube, three inches of which is bent downward from the horizontal to retain the fused salt. The residual gas is transferred to the receiver by the pump, brought to a constant temperature by a stream of hydrant water and measured in the usual manner. The mercury pump employed is a very effective one, of simple and easy construction, devised for use in this kind of analysis.

With regard to the handling of the soda-lime method the following results have been arrived at:—

1. Contrary to what is commonly stated, fine pulverisation of the substance to be analysed is not necessary. If the substance will pass holes of one millimetre in diameter it is fine enough.

A sample of dried blood which passed through a sieve with meshes one millimetre in diameter gave 7'58 per cent of nitrogen. A portion of the same, ground extremely fine with sand, gave 7'64 per cent.

Fish-scrap passed through the same sieve gave 8'98 per cent of nitrogen; when ground with sand, 8'95 per cent. A second sample of fish sifted as above gave 8'69 per cent nitrogen. By the absolute method it yielded 8'79 per cent.

2. Neither the highest heat possible to obtain in an Erlonmeyer gas combustion furnace, nor a long layer of strongly heated soda-lime, nor these two conditions united, occasion any appreciable dissociation of the ammonia formed in combustion.

A sample of dried blood gave 11'42 per cent of nitrogen when the combustion was made in a tube 14 inches long at a dull red heat.

The same sample yielded 11'56 per cent when deter-

* The use of a short layer of oxide of copper at the anterior end of the tube was adopted by the writer in 1873, when making analyses of Connecticut tobacco, he having found it impossible to make a blank combustion when using carbonic acid to sweep the tubes, without obtaining unabsorbed gas in the receiver, which proved to be combustible. By burning this gas in the analysis, he was able to get results on uric acid and potassium ferrocyanide agreeing closely with theory and with those obtained by the soda-lime method. More recently, Frankland has adopted this use of copper oxide in nitrogen estimations for water analysis.

Since the above note was printed I have learned that Finkener, in his edition of "Rose's Handbuch," anticipated me in this use of copper oxide.

S. W. J.

S. W. J.

mined in a tube 30 inches long, the mixture occupying 12 inches, and the rinsings and clear soda-lime 16 inches, using as high a heat as possible.

The same experiment was tried with egg-albumin. In a tube 14 inches long it yielded 12'25 per cent; in one 30 inches long, filled as above, 12'34 per cent.

A superphosphate containing animal matter yielded 3'02 per cent of nitrogen when the heat was kept quite low, 3'04 per cent when the heat during combustion was very bright red.

3. The use of pure sugar or of oxalic acid as a diluent does not in any way affect the result.

	Percent nitrogen.
Dried Blood, 0'5 grm., gave	10'33
" " with 0'5 grm. sugar gave ..	10'29
" " " 1'0 " " " ..	10'68*
Dried Blood " gave	11'56
" 0'37 grm. with 1 grm. sugar gave ..	11'53
Egg Albumin, 0'5 " gave	12'51
" 0'5 " with 0'5 grm. sugar gave ..	12'41
" 0'46 " " " " ..	12'50
" 0'3 " " " " ..	12'54

Experiments with oxalic acid gave similar results.

4. Iron tubes of proper length may be substituted for glass. The results are as satisfactory, but more time is required to make the combustion.

The iron tubes used in the following trials were 22 inches long. They were closed at the rear, and at the end of the combustion were cleared of ammonia by heating a mixture of oxalic acid or sugar and soda-lime, which was kept cool in the rear of the tube till needed for this purpose.

	Per cent nitrogen.	
	In glass.	In iron.
Egg Albumin	12'36 ..	12'25 ..
Dried Blood	11'42 ..	11'56 ..
Fish scrap.. ..	9'01 ..	8'99 ..
"	8'71 ..	8'68 ..
"	8'66 ..	8'72 ..
Ammoniated superphosphate..	2'69 ..	2'61 ..
"	2'71 ..	2'68 ..

In the Station Report for 1877 it was stated that combustions in iron tubes yielded 0'2 to 0'5 per cent less than those in glass. This deficiency was occasioned simply by using too short an anterior layer of soda-lime, the tubes being but 14 inches long. Since iron is a good heat conductor these tubes had an effective length much less than glass tubes of the same dimensions would have.

5. A suitable length of the anterior layer of soda-lime must be secured in order to get a good result. With 0'5 grm. of substances, such as are encountered in agricultural chemistry, containing less than 8 per cent of nitrogen, a glass tube of 12 to 14 inches is long enough. As the content of nitrogen increases to 10 per cent or over, we make the tubes several inches longer. In the combustion of dried blood or egg-albumin we prefer a tube 25 to 30 inches long, and the mixture of soda-lime and substance should occupy rather less than half the tube, a layer of pure soda-lime of 12 or more inches long being essential for perfectly destroying the volatile organic matters.

6. The long anterior layer of pure soda-lime must be brought to a full red heat before heating the mixture, and must be so kept throughout the combustion.

7. No fumes or tarry matters, indicative of incomplete combustion, should appear in bulb-tube or receiver.

8. When the combustion proper is begun under the conditions above described, it can be carried on quite rapidly until completed. The contents of the tubes then show no sign of unburned carbon.

9. We get equally good results whether the mixture is made intimately in a mortar or more roughly by stirring

* In this analysis the combustion was not complete, owing to the large amount of sugar being mixed with insufficient soda-lime.

with a spatula in a capsule or scoop, or by mixing in the tube with a wire.

10. We usually allow the glass tube to cool somewhat before aspirating with air to sweep out the ammonia, but have not as yet decided whether this precaution is essential.

11. We receive the ammonia of the combustion in a bulb-tube or flask containing standard hydrochloric acid, and we measure the excess of acid by a standard ammonia solution, using tincture of cochineal as the indicator.

In conclusion, we recall the fact that some careful experimenters—Ritthausen, Mærcker, Petersen, Kreusler—have repeatedly obtained results nearly as accordant as ours in the use of the two methods. Such results could scarcely be accidental, and we express our conviction that the discrepancies observed by others have been due to imperfect working of the processes. We have had great experience with both modes of determining nitrogen (one of us for twenty-five years), and not until recently have we learned how to be fairly certain of our results on all the classes of substances we have had occasion to analyse, either with the soda-lime process or by the absolute method. The purity and uniformly satisfactory qualities of the new soda-lime have greatly facilitated our work. The use of cochineal as an indicator we consider very favourable to exact titration.—*American Chemical Journal*.

NOTICES OF BOOKS.

An Introduction to the Practice of Commercial Organic Analysis; being a Treatise on the Properties, Proximate Analytical Examination, and Modes of Assaying the various Organic Chemicals of Preparations employed in the Arts, Manufactures, Medicine, &c., with concise Methods for the Detection and Determination of their Impurities, Adulterations, and Products of Decomposition. By ALFRED H. ALLEN, F.C.S., F.I.C., Lecturer on Chemistry at the Sheffield School of Medicine, Public Analyst of the West Riding of Yorkshire, Borough of Sheffield, &c. Vol. I. Cyanogen Compounds, Alcohols and their derivatives, Phenols, Acids, &c. London: J. and A. Churchill.

ON opening this book we were curious to learn how far it had been found practicable to carry out the promises implied by the above somewhat formidable title, and we feel bound to say that a scrutiny proved that the author had justified our anticipations. As stated in the preface, "it is a lamentable fact that while our young chemists are taught to execute ultimate organic analysis, and to ring the changes on the everlasting chloro-, bromo-, and nitro-derivatives of the bodies of the aromatic series, the course of instruction in our leading laboratories does not include even qualitative tests for such every-day substances as alcohol, chloroform, glycerin, carbolic acid, and quinine." Hence the author has set himself to work to compile a text-book of Proximate Organic Analysis, and has succeeded in producing a volume which is in a great measure unique, and which every practical chemist will find most useful. As explained in his preface, the author found the subject matter too extensive for compression into a single volume, and he has therefore reserved various important items for future publication, but the volume now issued is complete in itself and is often exhaustive on the analytical characters of the substances treated of. The author has divided the volume into seven parts, the first of which, extending over 20 pages, is introductory. This division, in our opinion, is the weak point of the book, and might have been wholly omitted without much loss. However, it contains a useful table of the physical properties and solubilities of a number of

organic bodies, and another table of the specific rotatory power of organic liquids. The second division treats of "Cyanogen and its Derivatives," and extends over 43 pages. All the well-known methods of assaying hydrocyanic acid, cyanides, ferrocyanides, &c., are well and clearly explained, and not a little novel matter is incorporated. Indeed, the amount of original information is one of the chief features of the book, there being few of the principal articles which do not contain methods or observations which are the personal experience of the author.

The third division, extending over some 70 pages, treats of the "Alcohols," but the only members of the series fully dealt with are methyl, ethyl, and amyl alcohols, and glycerin. Under ethyl alcohol there is a full description of the methods of examining cider, spirits, liqueurs, and tinctures, but wine and beer are not treated at length in this volume. Methods for the determination of nearly all their chief constituents are, however, to be found scattered throughout the book.

In the next division Mr. Allen treats of "Neutral Alcoholic Derivatives," under which description he classes Compound Ethers, Chloroform, Chloral, &c. As an illustration of the completeness of the book we may say that, as sub-articles under the head of chloroform, the author describes and gives modes of assaying bromoform, iodoform, and methene dichloride, while under "Chloral," butyric or croton chloral, dichloride of propylene, and chloroacetic acid are treated of. As a rule the author limits his description to such of the properties of the bodies as are of analytical interest, and the modes of preparation are in most cases very meagrely described.

In the remaining divisions the author treats of "Acid Alcoholic Derivatives and Vegetable Acids," "Phenols," and "Acid Derivatives of Phenols." Space will not allow us to notice these divisions at greater length, but the whole of the information appears to have been carefully compiled, and brought up to the latest date. The author does not confine himself to a description of the reactions and methods of determining organic bodies in a state of purity, but gives practical directions for the assay of such products as tartars, lime and lemon juices, preparations of carbolic acid, tanning materials, and oil of bitter almonds. Mr. Allen assumes the possession of a fair amount of skill in manipulation, and hence the work is well adapted for use by professional and works-chemists, public analysts, and advanced students.

On page 324 the formula of picric acid is written $C_6H_2(NO_2)OH$, but it is shown with three atoms of NO_2 on a subsequent page. On page 326 we are told that "phenyl-sulphuric is a decided antiseptic," the word "acid" being omitted. A formula, said in the list of *errata* to be found on page 176, really appears on page 166. With the exception of these, and a few similar errors of minor importance, the book is remarkably free from mistakes, and is printed in clear type.

We congratulate Mr. Allen on the way he has accomplished a very difficult task, and shall look forward to the issue of the second volume of his original and very useful work.

The Elements of Systematic Qualitative Analysis. By J. H. SNIVELY. Nashville, U.S., 1878.

Tables for Systematic Qualitative Analysis. Same Author.

WE have vainly searched for any special feature that we could commend in Mr. Snively's two small books. It is impossible that they should meet with acceptance in England, where manuals of qualitative analysis, at once more exact, more complete, and more moderate in price, are in common use. The author claims no originality, save in arrangement of materials, acknowledging his obligations to Fresenius, Galloway, Attfield, and other writers and workers on the methods of analysis. But the 170 pages to which his manual extends are so incomplete

and unsatisfactory that we cannot imagine the authorities he has consulted feeling gratified at the help they have afforded Mr. Snively. The 33 woodcuts with which the book is disfigured are beneath criticism, and can but provoke a smile. Note especially the elegant form of the agate mortar on page 6, and the marvellous perspective of the test-tube rack on page 23. It is scarcely worth while to dwell any further upon the contents of Mr. Snively's manual and tables, but anyone may test the justness of our criticism by reading the section on "Reagents," pp. 31 to 46, noting particularly the directions for preparing lime-water, gold chloride, chlorine-water, platinum tetrachloride, and potassium nitrite.

Twenty Lessons in Inorganic Chemistry. By W. G. VALENTIN. W. Collins, Sons, and Co., 1879.

THIS book is one of the most recent of the countless books and booklets to which the Science and Art Department has given birth, and it is undoubtedly one of the very best of the extensive group of works to which it belongs. The late Principal Assistant in the Royal College of Chemistry, South Kensington, Mr. W. G. Valentin, was a most assiduous and energetic teacher, an ingenious and apt manipulator, and a sound chemist. These twenty lessons on the non-metals exhibit in every direction the peculiar merits of their author. He has wisely abandoned the complex formulæ of Dr. Frankland, although explaining the system on which they are constructed. Moreover, he has interspersed his several chapters with a host of little bits of useful information, hints as to the successful performance of experiments, and facts and data not readily accessible elsewhere, so that the volume, small as it is, and elementary as it is meant to be, is not at all like its many rivals in the same field. The illustrations, likewise, are unusually excellent, representing really practicable forms of apparatus, and experiments that can be performed. Although the lines followed by Mr. Valentin are those traced out by the "Science Directory" for students of the elementary grade, we feel sure that the volume before us will be found of much greater value and of much wider usefulness than the ordinary "cram and sham" books to which so many examinees have resort.

CORRESPONDENCE.

VOLUMETRIC DETERMINATION OF CHROMIUM.

To the Editor of the Chemical News.

SIR,—Mr. Galbraith, in a letter to the *CHEMICAL NEWS* (vol. xxxix., p. 276), refers to a previous publication by himself of a method of estimating chromium, similar to that described by me in the *Journal of the Chemical Society*. Mr. Galbraith's communication had escaped my notice. His method is the same in principle as mine, and I have been unfortunate in having failed to notice it sooner. I am, however, not without hope, that the improved detail and extended application of this method, as shown in my paper, may be of use to those chemists interested in the analysis of chrome iron-ore.—I am, &c.,

W. J. SELL.

July 8, 1879.

SULPHUR IN ICELAND.

To the Editor of the Chemical News.

SIR,—In your review of Dr. Lunge's "Treatise on the Manufacture of Sulphuric Acid and Alkali" (*CHEMICAL NEWS*, vol. xxxix., p. 287), the statement is made that "the sulphur deposits of Iceland and Saba remain undeveloped." This is incorrect so far as regards Iceland, the fact being that the working of the deposits in Guldbringa Syssel in the south-western corner of the island is being actively prosecuted with satisfactory results.—I am, &c.,

THOS. G. PATERSON.

53, George Street, Edinburgh, July 9, 1879.

University of London.—The following candidates have passed the recent D.Sc. Examination:—Branch VI. *Electricity*—(a) *Treated Mathematically*. Hugh William McCann, Trinity College, Cambridge. (b) *Treated Experimentally*. John Ambrose Fleming, St. John's College, Cambridge. Branch XII. *Vegetable Physiology*. Sydney Howard Vines, Christ's College, Cambridge.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

JUNE, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

	Appearance in a foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia.	Chlorine.	An- Sulphuric hydride.	Hardness on Clark's Scale.		
		Saline.	Organic.								Before Boiling.	After Boiling.	
													Grains.
<i>Thames Water Companies.</i>													
Grand Junction	Slightly turbid	0'000	0'012	0'129	0'165	21'00	6'940	0'570	1'150	1'53	13'0	4'2	
West Middlesex	Clear	0'000	0'012	0'135	0'037	20'30	7'280	0'720	1'150	1'53	12'6	3'7	
Southwark and Vauxhall	Clear	0'000	0'012	0'102	0'129	19'11	6'660	0'500	1'000	1'83	12'1	3'3	
Chelsea	Clear	0'000	0'012	0'120	0'033	19'80	6'550	0'790	1'220	2'06	12'0	3'3	
Lambeth	Clear	0'000	0'012	0'120	0'070	20'20	6'160	0'570	1'160	2'16	12'6	5'1	
<i>Other Companies.</i>													
Kent	Clear	0'000	0'003	0'450	0'002	28'10	8'960	0'910	1'580	2'80	17'6	8'5	
New River	Clear	0'000	0'006	0'120	0'059	20'60	7'840	0'800	1'150	1'80	13'2	3'7	
East London	Clear	0'000	0'011	0'150	0'048	21'00	8'060	0'460	1'150	1'66	13'2	4'2	

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 25, June 23, 1879.

Absorption of Ultra-violet Rays by the Atmosphere.—A. Cornu.—The author brings forward arguments, which, if not absolutely demonstrative, render it extremely probable that the real extent of the solar spectrum on the more refrangible side is considerable, and that it is abruptly cut off by the intervention of a foreign cause. This cause he shows experimentally to be atmospheric absorption. It remains to be ascertained what respective parts are played by the constituents of the atmosphere in this absorption.

Inexact Application of a Dynamical Theorem made by MM. Bertin and Garbe to explain the Movements of the Wings of the Radiometer.—A. Leduc. (See *Comptes Rendus*, lxxiv., p. 30).—It results from the objections raised by the author that the cause of the movement of the radiometer must still remain in reserve. The foregoing considerations take no account of the nature of the fluid inclosed in the globe. The reservation of judgment here demanded is corroborated by the diversity of the theories advanced by the most distinguished physicists of all countries to explain the movements of the discs. It is plausible to believe that the cause sought for must be complex, and that we have a superposition of several effects. Attempts should be made to particularise these effects according to a programme of systematised experiments, which may be thus summed up:—Varying the substance of each disc and of its surfaces according to the diverse thermic and luminous properties of the materials employed; modifying successively the nature of the luminous rays and polarising them in various directions with respect to the discs; suspending the globe on two very fine points and enclosing it in a receiver, which is then exhausted.

Representation of the Solar Spectrum.—M. Thollon.—The design laid before the Academy was executed in Italy. It is 10 metres in length from A to H, and is composed of about 4000 rays. For the present he merely calls attention to the singular resemblance of the groups A and B, which have not hitherto been resolved in as complete a manner, and proposes a novel classification of the solar rays according as they are formed of a nebulosity without nucleus, of a nucleus without nebulosity, of a nucleus and a nebulosity, the latter predominating, or of a nucleus and a nebulosity, the nucleus predominating.

Study of the Molecular Constitution of Liquids by means of their Coefficient of Expansion, their Specific Heat, and their Atomic Weight.—R. Pictet.—Not susceptible of abstraction.

Alloys of Lead and Antimony and the Liquations and Supersaturations which they Present.—F. de Jussieu.—Noticed elsewhere.

Production of Hydrocellulose.—A. Girard.—The definite compound which the author has made known under the name of hydrocellulose, and which was obtained by submitting cellulose to the action of liquid or dissolved acids may likewise be formed by the action of gaseous acids.

Reversion of Superphosphates.—H. Joulie.—Superphosphates, even when heavily charged with iron and alumina, if prepared with a sufficient quantity of acid do not undergo a reversion of the assimilable phosphoric acid (soluble in basic ammonium citrate), but they are apt to remain in the state of a soft, elastic paste, unfit for distribution in the soil. If the dose of acid is reduced

and the action is incomplete, the mass dries better, but the assimilable phosphoric acid undergoes reversion by the action of the sesquioxides upon the mono- and bicalcic phosphates formed at first, the result being iron and aluminium phosphates, along with tricalcic phosphate, much less soluble in ammoniac citrate than the compounds first mentioned. The addition to superphosphates of chalk, or of gypsum containing carbonate of lime as dryers, produces immediately the same phenomenon, which increases with time.

Presence of Oxygen in the Sun.—H. Draper.—The author submitted to the Academy a photographic proof of the blue and violet part of the solar spectrum, and of the spectrum of oxygen. The coincidence of the brilliant rays of oxygen with the brilliant bands of the solar spectrum is a proof in favour of the existence of oxygen in the sun.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 2, 1879.

Anthracen Compounds belonging to the Chrysazin Series.—C. Liebermann.—The author describes the di-sulph-anthracenates of sodium, potassium, calcium, and barium, chrysazol, diacetyl-chrysazol, diacetyl-chrysazin, chrysazin, tetra-nitro-chrysazin potassium, and the corresponding magnesium compound, also tetra-nitro-anthra-rufin potassium and sodium.

The Reduction of Sulph-anthraquinonic Acids.—C. Liebermann.—The author, on treating the salts of mono-sulph-anthraquinonic acid with hydriodic acid (sp. gr. 1.7) and red phosphorus for some hours with an ascending condenser, obtains the corresponding mono-sulph-hydride-anthracenates.

Action of Potassic Carbonate upon Iso-butyl-aldehyd.—F. Urech.—The resulting compound is a semi-fluid, sparingly soluble body, which on distillation is resolved into iso-butyl-aldehyd and certain condensation-products which the author is examining in detail.

Limits of the Applicability of the Method of Determining Vapour-densities in the Barometric Vacuum.—J. W. Brühl.—This paper cannot be usefully abstracted without the accompanying engravings.

A Process for the Purification of Mercury.—J. W. Brühl.—The mercury is shaken up with an equal volume of a solution of 5 grms. potassic bichromate in 1 litre water with the addition of a few c.c. of sulphuric acid, the operation being continued till the water becomes of a pure green. A strong current of water is then introduced into the flask, which washes away a green powder. This process is repeated if needful, and the mercury is finally agitated with distilled water till its surface remains perfectly bright.

No. 3, 1879.

Formation of Sulphonic Acids from Sulphones.—R. Otto.—If two molecules of mono-chlor-hydrin-sulphuric acid are allowed to act upon 1 mol. of the sulphon, the disulphonic acid of sulpho-benzid is formed.

Preparation of Sulphuretted Hydrogen in Chemical Investigations.—R. Otto.—The author maintains that if either the iron-sulphide or the acid employed is contaminated with arsenic arseniuretted hydrogen is evolved. It is better to prepare the gas by the action of hydrochloric acid upon calcium or barium sulphide, as recommended by Mohr and Grothe. Myers suggests that the traces of arsenic which have been found in the normal animal body may perhaps be due to impure hydrogen sulphide.

Action of Iodine upon Aromatic Compounds with long Lateral Chains.—K. Preis and B. Raymann.—Not suitable for abstraction.

Contributions to a Knowledge of Cholesterolin.—K. Preis and B. Raymann.—The authors describe the action

of fuming nitric acid upon cholesterin and upon cholesteryl-chloride.

Certain Azo Compounds.—P. Weselsky and R. Benedikt.—An account of azo-benzol-phloro-glucin, para-azotoluol-phloro-glucin, para-azo-phenol-phloro-glucin, and the azo derivatives of naphthylamin.

Oxidation-products of the Cinchona Bases.—Z. H. Skraup.—This paper does not admit of useful abstraction.

Contribution to a Knowledge of Ultramarine.—E. W. Büchner.—By heating sodium, aluminium, and silicium in a current of sulphuretted hydrogen the author obtained a black mass, which, on extraction with water and re-heating with free exposure to the air, became converted into ultramarine-blue.

Constitution of Phenanthren.—G. Schultz.—The author considers he has proved that diamido-diphenic acid from diphenic acid is identical with that obtained from meta-nitro-benzoic acid.

Phthalein of Ortho-cresol.—G. Fraude.—Not adapted for useful abstraction.

Contribution to the Knowledge of Glyoxylic Acid.—C. Böttinger.—An account of the action of ammonia upon glyoxylic acid.

The Constituents of the Etheral Oils of Certain Ericaceæ.—H. Köhler.—An examination of the oils of *Gaultheria punctata* and *G. leucocarpa*.

The Heat Liberated on the Contact of Anhydrous Sodium Sulphate and Water.—L. C. de Coppet.—The author rejects Thomsen's explanation of the rise of temperature as a consequence of the formation of the hydrate, $\text{Na}_2\text{SO}_4\cdot\text{H}_2\text{O}$.

Specific Rotatory Power of Iso-cholesterin.—E. Schulze.—Of the two isomeric fatty bodies of wool iso-cholesterin is dextro-rotatory, whilst cholesterin is lævoro-rotatory.

Determination of the Specific Weight of Pulverulent Bodies.—F. Rüdorff.—The method proposed cannot be intelligibly described without the aid of the accompanying engraving.

Contributions to a Knowledge of Methyl-crotonic Acid and of Angelic Acid.—E. Schmidt.—The conversion of angelic acid into methyl-crotonic acid takes place spontaneously in course of time, and in the behaviour of the former a certain resemblance to β -crotonic acid cannot be overlooked.

Azo-derivatives of Diphenyl-amin and Diphenyl-nitrosamin.—O. N. Witt.—An account of the "tropeolin" colours, a name which the author applies to all yellow and orange dyes produced by introducing the sulpho group into amido or oxyazo bodies. The most beautiful among these is the potassium salt of phenyl-amido-azo-benzol-sulphonic acid (tropeolin OO) which gives a fiery golden yellow on silk and wool. The basic properties of amido-azo-benzol are so far neutralised that its application is possible in presence even of weak acids.

A New Universal Support for the Pocket Spectroscope.—F. v. Lepel.—Unintelligible without the accompanying engravings.

The Chemistry of Chondrin.—R. Petri.—On treating chondrin with sulphuric acid and steam the author obtains bodies resembling syntonin, peptons, and a crystalline nitrogenous substance, free both from albumen and peptons. It has the character of a polybasic acid.

Chemiker Zeitung.
No. 20, May 15, 1879.

The proposed duty of 100 francs per 100 kilos. on the importation of patent and proprietary medicines is denounced as implying an official recognition of such articles

(a well-known result of the stamp-duty on patent medicines in England).

Prof. Knop denounces "dilettantism in sanitary chemistry," and accuses Dr. Elsner of neglecting analytical precautions, the necessity of which he had formally admitted.

Separation of Ferric Oxide and Alumina from Manganese.—A. Classen's method is based upon the fact that solutions of manganic salts are precipitated by neutral potassium oxalate, the precipitate being soluble in an excess of the latter. The potassium-manganic oxalate is then decomposed by concentrated acetic acid, in which manganese oxalate is insoluble, whilst potassium ferric oxalate formed in the same manner gives no precipitation with acetic acid. To obviate the disturbing action of the alkaline chlorides which prevent the complete precipitation of the manganese oxalate, Classen adds zinc chloride in excess, so that the ZnO may be three or four times the quantity of the MnO . All the manganese is thus precipitated with the zinc oxalate, the cold bulky precipitate is rendered dense and crystalline by heating to 40° to 50° , and is then washed, dried, and ignited with excess of air, forming a mixture of manganese and zinc oxides which often, in consequence of imperfect washing, contains traces of potassium permanganate. The contents of the crucible, after a short ignition, are digested with alcohol and hot water, and washed again. The ignited mixture of Mn_2O_3 and ZnO is decomposed by concentrated hydrochloric acid, the chlorine set free is passed into solution of potassium iodide, and the iodine which separates is titrated with sodium hyposulphite. The filtrate from the oxalates contains ferric oxide and alumina, which, after concentration, are precipitated with alcohol and a little acetic acid, and the ignited precipitate is washed with water containing a little potassium carbonate. The method is likewise suitable for separating calcium, cobalt, nickel, and probably copper from ferric oxide and alumina.—*Zeit. Anal. Chemie*, 1879, p. 175.)

Separation of Manganese and Zinc.—Tamm (Guyard?) proposed to determine manganese as carbonate by carbonate of ammonium, ammonium chloride having first been added, and applies this reaction to the separation of manganese and zinc. A. Classen points out that the separation is very imperfect.—*Zeit. Anal. Chemie*, 1879, 194.

Volumetric Determination of Free Oxygen in Gaseous Mixtures by Means of Phosphorus.—To determine oxygen in the chamber-gas of sulphuric acid works, Lindemann, after removing the nitrous acid by means of concentrated sulphuric acid, uses, in place of sodium pyrogallate, moist phosphorus, which he introduces in the shape of exceedingly thin rods into the absorption-vessel of an Orsat's apparatus. The traces of phosphorous acid not absorbed by the water have no influence upon the result on account of their slight tension.—*Zeit. Anal. Chemie*, 1879, 158.

Volumetric Determination of Zinc.—C. Mann.—Hydrous zinc sulphide and silver chloride are readily and completely transformed into silver sulphide and zinc chloride. The amount of chlorine present is then ascertained by Volhard's titration process with silver, and the zinc is thus calculated.—*Zeit. Anal. Chemie*, 1879, 162.

Separation of Arsenic and Antimony.—Bunsen's process for separating these two metals by prolonged digestion with potassium sulphide and excess of sulphurous acid has been re-examined by Nilson and found untrustworthy. Bunsen no longer employs this method.—*Zeit. Anal. Chemie*, 1879, 165.

Cupreous Ammonia.—A. Schalm obtained a cupreous ammonia, perfectly colourless, but which gave a black-brown copper sulphide with sulphuretted hydrogen. The copper is present as cupro-cyan-ammonium.—*Zeit. Oest. Apoth. Ver.*, 17, 189.

No. 21, 1879.

This issue contains nothing calculated to interest our readers.

No. 22, 1879.

The Anglo-Swiss Condensed Milk Co., in reply to the accusations of Dr. Soxhlet and of Baron H. v. Liebig, produce numerous analyses of their condensed milk from Profs. Schulze, Mærcker, Schwarzenbach, Wagner, and others, showing that the cream is not removed before or during the process of condensation.

A specimen of liquid sulphurous acid on view at the Berlin Exhibition exploded with some violence. Fortunately, no one was hurt.

Determination of Superphosphates.—H. Albert and L. Siegfried.—The authors maintain that the hydrated basic and neutral phosphates are substantially equal in value to phosphoric acid soluble in water. A determination of the latter alone does not give the agricultural value of a superphosphate, and is irrational, as it excludes from use phosphorites and other aluminiferous phosphatic minerals. A determination of the total phosphoric acid soluble in ammonium citrate is declared preferable.—*Zeit. Anal. Chemie*, 1879, 220.

Determination of Phosphoric Acid in Fish-guano.—B. E. Dietzell and M. G. Kressner.—In analyses of "Polar" and "Lofoden" guanos the authors obtained concordant results by a twice-repeated evaporation of the ash with concentrated nitric acid. Fusion with soda nitrate in case of Lofoden guano yielded too high results, a certain portion of phosphorus being present not as phosphate, but in organic compound.—*Zeit. Anal. Chemie*, 1879, 225.

Detection of Foreign Fats in Butter.—J. Koettstorfer.—The author, taking advantage of the fact that butter contains more acids of low molecular weight than other fats, and consequently requires more potassa for saponification, places from 1 to 2 grms. of the filtered sample in a tall covered beaker, holding about 70 c.c., along with 25 c.c. of an alcoholic solution of potassa of about semi-normal strength, and saponifies by heating for fifteen minutes in the water-bath. He then titrates with semi-normal hydrochloric acid, using a very dilute alcoholic solution of phenol-phthalein as indicator. The standard of the potassa solution not being quite permanent is re-determined each time with 25 c.c. heated in the same manner. 1 gm. pure butter fat requires for saponification from 221.4 to 232.4 milligrms. KHO; 1 gm. mutton tallow, 197 ditto; beef tallow, 196.5; hogs' lard and oleo-margarin, 195.5; olive oil, 191.8; and rape oil, 178.7. The author considers his process available for the quantitative detection of rape oil in olive oil.—*Zeit. Anal. Chemie*, 1879, 199.

According to Gawalowski the filter-papers of A. Schmidt, of Gross Üllersdorf, in Austria, are preferable to the Swedish "Munktell." They are much cheaper, very equal in texture, and yield a smaller proportion of ash.

Justus Liebig's Annalen der Chemie,
Band 196, Heft 2.

Nicotin and Nicotinic Acid.—R. Laiblin.—The author gives a method for obtaining pure nicotin in quantity, and then proceeds to the examination of nicotinic acid, $C_6H_5NO_2$, and its derivatives. He considers this acid as pyridincarbonic acid, and attempts its synthesis, but unsuccessfully.

Nature of Cohesion and its Chemical Significance.—Fried. Mohr.

On Chloranilins and Chloronitrilins.—F. Beilstein. and A. Kurbatow.—An extensive memoir, not capable of useful abstraction.

Experimental Researches on Hydrogen Peroxide.—Em. Schoene.—(Fifth Memoir: Behaviour of Hydrogen

Peroxide with Ozone and Chlorine).—In the mutual decomposition of ozone and hydrogen peroxide both lose equal quantities of oxygen, whilst the gas in which the former is present expands by a volume equal to the sum of the volumes of oxygen lost by both bodies. The quantity of oxygen evolved in the reaction of chlorine and hydrogen peroxide is double the quantity which the latter contains in excess above water.

Chlorides of Camphor.—F. V. Spitzer.—The author has prepared these compounds by treating camphor with phosphorus chloride.

Les Mondes, Revue Hebdomadaire des Sciences,
No. 6, June 5, 1879.

A new application of the electric light is proposed by M. Trouvé. It is a brilliant light in a closed vessel by means of which it is possible to explore the most "fiery" mines without danger.

M. L. Digeon considers that the phylloxera is most to be dreaded in its winged state, as it is then easily conveyed by the winds. He proposes to syringe the vines from time to time with milk of hydraulic lime.

Nos. 7 and 8, June 12 and 19, 1879.

These issues contain no original chemical matter.

Reimann's Fürber Zeitung,
Nos. 18 and 19, 1879.

The most important article in these two issues is a paper by Dr. A. Müller-Jacobs "On the Use of Tannin in Turkey-red Dyeing on the New Principle." The author says:—"The whole art of dyeing, and especially Turkey-red dyeing, depends essentially on a dialysing process, and the object of the oiling or mordanting is merely to convert the vegetable fibre into the best possible dialyser. Parchment (paper?) and mercerised cotton are dialysers, and both possess in a high degree the property of taking up without mordant such colours as dye without base. The white-mordanted tissue (oiled on the old principle) serves as a dialyser for solutions of alumina. Simultaneously, with the dissociation of the aluminium salts the hydrate of alumina combines with the fixed oleic acid. Of course in this process the chemical affinity of the fixed oil for aluminic salts cannot be overlooked. Mercerised cotton likewise decomposes salts of alumina, and this is still more completely effected by cellulose first mercerised and then oiled on the new method. The process in the dye-bath depends still more upon dialysis than does the mordanting process. The most fiery red is obtained by converting the goods into the best possible dialyser. The chemical union of the alizarin with the mordants occurs only at the more elevated temperatures of the dye-beck, when the whole of the colouring matter has been absorbed and the bath has become almost colourless. The more readily the cotton is dialysed by the mordanted cotton (which occurs at low temperatures), i.e., the better the colouring matter 'works on' at a low temperature, the more excellent is the red. Additions of chalk, of aluminous salts, &c., even in the smallest quantity, alter essentially the liquid to be dialysed, and have thus a powerful influence on the colour to be produced. This seems to show that the process of dyeing is physical rather than chemical in its nature. On the addition of small quantities of chalk to the dye-bath, e.g., $\frac{1}{2}$ lb. to 100 lbs. goods, the colouring matter works on much more quickly, the dialysis being accelerated; whilst, on the other hand, the addition of Turkey-red oil, of acids, or alkalies, even in the smallest quantities, or of certain salts, prolongs, or even prevents the dialysis. Cotton, if possible, slightly mercerised and treated with Turkey-red oil, receives by the aid of tannin the power to dissociate

aluminic compounds as in the old process, and to fix the hydrate upon the fibre in the state best suited for the production of brilliant colours. By the addition of tannin we compensate for various defects relating to dialysis, and render the addition of Turkey-red oil to the dye-bath possible, whereby the alizarin is absorbed more slowly and a greater equality in the colour is obtained. By fixing large quantities of alumina in the process of mordanting we produce a very fiery red, which lies more on the surface, whilst the fibre remains white internally and the colour scarcely bears a strong soaping. This evil may be avoided by adding tannin to the dye-bath along with Turkey-red oil, thus retarding the dialysis and giving the colour time and opportunity to penetrate into the interior of the fibre. Without tannin we should produce a tawny fugitive red; the addition of chalk (without oil and tannin) would give a red which would not bear clearing. An excess of tannin gives shades which are very fast, but brownish.

No. 20, 1879.

This issue contains nothing of general interest.

No. 21, 1879.

This issue is chiefly taken up with a notice of the Berlin Industrial Exhibition, and with extracts from M. Moyret's exposure of the process of weighting silks.

Behring's Thickener (Glutine).—This adhesive matter consists of casein dissolved in sodium tungstate. It is recommended for a variety of purposes, but for thickening printers' colour it can only be used where the reaction of tungstic acid with the dye-warens present would produce no objectionable effects.

No. 22, 1879.

M. Moyret's paper on weighted silks is continued. For whites and bright shades sugar is used, to which the author proposes an addition of a decoction of quassia. Silk dresses thus prepared would serve for fly-papers. Stannic chloride, mysteriously spoken of in the trade as X, is employed in addition to sugar. Barium sulphate has also been proposed, but it deprives the silk of its lustre.

No. 23, 1879.

Malachite Green and Benzol Green (Victoria Green).—The writer points out, with reference to these two colours—probably identical—a curious discrepancy between the patent system of France and that of Germany. In the former country a chemical product, as such, however produced, may be the subject of a patent. In Germany the process of preparation only is patentable. Thus, if malachite green and Victoria green are identical, the latter colour cannot be made or sold in France except under Dœbner's patent. The author mentions this as an instance showing the necessity for an international patent law.

No. 24, 1879.

According to M. Moyret, 40,000 kilos. of copperas, pyrolignite of iron, and other iron mordants are used daily in the Lyon district for weighting black silks.

A CHEMICAL REMINISCENCE.

There are huge heaps of tank-waste by Sankey's dull stream,

Hydric Sulphide is reeking there all the day long;
In the hours of my childhood 'twas like a bad dream

To stay o'er the tank-waste and sniff the fumes strong.

That stream with its odours I ne'er shall forget,

And often when musing on days that are o'er

I think if the tank-waste is lying there yet

Are the vapours still choking on Sankey's grim shore?
S.

MISCELLANEOUS.

Reform Club.—Mr. Chas. W. Vincent, F.R.S.E., F.C.S., has been appointed the colleague and successor of Mr. Henry Campkin as Librarian of the Club.

Cerium and Didymium.—Dr. Theodor Schuchardt, of Goerlitz, in a letter to the Editor in May last, stated that he had succeeded in obtaining the metal cerium by electrolysis in globules weighing from 4 to 5 grms. Under date of July 12, Dr. Schuchardt says he has just succeeded in obtaining from didymium chloride, by electrolysis with six Bunsen elements, the metal didymium in globules the size of a small pea.

Valentin Memorial Fund.—We learn with pleasure that many of the friends and former pupils of this well-known Chemist have repented to the appeal, which as we stated some weeks since, has been made on behalf of the family. There must, however, be others whom previous applications have not reached; we would therefore mention that subscriptions will still be welcomed by the Hon. Treasurer of the Fund, Mr. F. W. Bayley, Royal Mint, E.

Royal Institution of Great Britain.—At the General Monthly Meeting, on Monday, July 7, 1879, Dr. C. W. Siemens Vice-President, in the chair, Messrs. Francis Fesser, James Garnett Heywood, Edmund de Quincey Quincey, Hal Smith, Herbert A. Taylor, and Miss Henrietta Lambert, were elected Members of the Royal Institution. The Special Thanks of the Members were given to the Earl Bathurst, M.R.I., for his munificent Present of a large Bust of William Hyde Wollaston, M.D., F.R.S., M.R.I., by Chantrey.

Royal School of Mines.—The following students have passed the Examinations which entitle them to the distinction of Associate of the Royal School of Mines:—In the Mining and Metallurgical Divisions—Mirza Mehdy Khan. In the Mining Division—W. E. Benton, A. G. Charleton, A. D. Ellis, E. N. Fell, H. B. Slatter, H. Strickland. In the Metallurgical Division—W. B. M. Davidson, A. H. Fison, F. W. Grey, E. Halse, E. W. Harvey, Malcolm Hill, J. H. Lucas, Walter March, A. Gordon Salamon. The two Royal Scholarships of £15 each have been awarded to J. J. Hood and J. F. Wilkinson. The Royal Scholarship of £25 has been awarded to Ralph G. Scott. The De la Beche Medal for Mining to A. D. Ellis, and the Murchison Medal for Geology to B. Mott.

NOTES AND QUERIES.

Soap.—Where can I get information to bleach the oil extracted from olive cake by CS₂ so as to produce a pure white soda soap? The soap I now manufacture has a brownish colour if from old, and greenish if from new oil—R. H.

Wash-bottles.—Several plans have appeared lately in the CHEMICAL NEWS showing appliances for shutting off communication between the interior of wash-bottles and the atmosphere, when it is desired to keep up a jet of water. In this connection allow me to suggest that a small hole bored through the cork is the best and simplest contrivance. By placing the forefinger over the hole it may be perfectly closed without any trouble.—P. CASAMAJOR.

Tetrachloride of Carbon. (Reply to C.I.C.)—Your correspondent, C.I.C., may not have seen Bloxam's account of the method of making this substance. Chlorine dried by passing through pumice wetted with strong sulphuric acid, is then made to traverse a Wolff's bottle containing bisulphide of carbon, and is carried onwards through a porcelain tube wrapped in sheet-copper, and filled with fragments of broken porcelain. Kept at a red-heat by a charcoal- or gas-furnace. The products are then condensed in a bottle surrounded by ice. These products when shaken with solution of potash are decomposed. The tetrachloride of carbon then separates and falls to the bottom. The upper layer having been poured off, the tetrachloride may be purified by distillation. (Bloxam's "Chemistry," Churchill, 1875; page 167; see diagram.) I should myself be glad to know of any better and more convenient process than the above.—F.C.S.

TO CORRESPONDENTS.

G. M. W.—To give the information you require would necessitate writing an essay on the subject.

J. C. H.—Phenic acid or carbolic acid.

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THE CHEMICAL NEWS.

VOL. XL. No. 1026

ON MANURE PHOSPHATES.

By K. WALTER,
Chemical Engineer, Aurelais, Belgium.

ALL natural and manufactured manure phosphates may be divided into the following four groups:—

- (1.) Phosphates which are only soluble in acids, and not, or only very slightly, soluble in oxalate of ammonia.
- (2.) Phosphates which are quite or for the greater part soluble in the oxalate.
- (3.) Phosphates which are quite or for the greater part soluble in citrate of ammonia; and
- (4.) Phosphates which are quite or for the greater part soluble in water.

The value, however, of phosphates for agricultural purposes cannot strictly be classed in these four groups. There are soils in which a given quantity of a natural phosphate of, say, 20 per cent of phosphoric acid, has the same effect as the same weight of a superphosphate of 12 to 14 per cent of phosphoric acid. But the price of the first would be about £1 15s. or less, while the price of the latter is at least £4 per ton. On the contrary, there are other soils which absolutely require superphosphate or precipitated phosphates. In some parts of France, mostly in Normandy, for instance, natural phosphates are nearly exclusively used for manuring purposes, and this with excellent effect. The money values of the different manure phosphates for different soils cannot be put into strictly divided classes. In the excellent works of H. Toulie* the subject of chemical manuring is extensively treated.

To the first of the above-named groups belong chiefly the richer phosphates, such as those of Canada, Curaçao, Navassa, Caçeres, and some other Spanish apatites; then the phosphates of Cambridge, Nassau, Bavaria, Norway, some kinds of the phosphates of the French Ardennes, Lot, Cher; and, finally, the Russian phosphates.

Those phosphates, as well as bone-ash, in which 25 per cent of their phosphoric acid is soluble in oxalate of ammonia, cannot be used with advantage for agricultural purposes, even when finely pulverised. They are nearly exclusively employed for the manufacture of superphosphates. For the manufacture of precipitated phosphate they are much too expensive; natural phosphates containing about 35 to 40 per cent of phosphate of lime—too poor for superphosphate making—being alone profitable.

To the second group belong those precipitated phosphates which have been simply precipitated with milk of lime without observing the necessary precautions. They contain for the most part tribasic phosphate of lime with very little dibasic of the same. This article is not specially manufactured, and only appears in the market as by-product of glue works. It is chiefly produced in France, and the present value in the Paris market is £6 per ton with 30 per cent of phosphoric acid. This phosphate contains on an average 27 to 31 per cent of phosphoric acid, and is used successfully in France for the manufacture of artificial guanos, also for the manufacture of superphosphate. It is but little employed directly for manuring.

* "Guide pour l'achat et l'emploi des Engrais Chimiques," par H. Toulie, Pharmacien en chef de la maison municipale de santé Administrateur délégué de la Société anonyme des produits chimiques agricoles. Cinquième édition; Société anonyme, 10, Quai de la Marne à Paris la Villette, 1876. "Méthode citro-urannique pour le dosage de l'acide phosphorique dans les phosphates et les engrais," par H. Toulie, Extrait Complémentaire du *Moniteur Scientifique Quésneville*. Paris, au Bureau de l'Annuaire, 12, Rue de Buci; 1876.

If this product is made without a surplus of lime, and dried at a temperature not exceeding 100° C. (=212° F.), it is found, after long experience, to have the same effect on soils which do not contain too little humus. This property has hitherto been recognised by few chemists. However, the time will come when these phosphates will be judged according to their value.

In a well manufactured product of that kind, 90 to 97 per cent of the contained phosphoric acid are soluble in the oxalate, and of those again 25 to 35 per cent in the citrate of ammonia. If, however, this precipitated phosphate is made with a surplus of lime, or dried at too high a temperature, it loses its property of being assimilated by the soil as quickly as superphosphate, and becomes as soluble in citrate as in oxalate.

Next there belongs to this second class the phosphate in the different guanos which contains 50 to 85 per cent of their phosphoric acid soluble in the oxalate. The phosphoric acid in the guano, though only very slightly soluble in the citrate, and in water only in traces, always was considered as highly assimilable. To treat the guano with sulphuric acid, as it became the fashion on the Continent some years ago, is merely done to fix the ammonia.

Further on we have to put in this category the phosphate of the raw bones and of the black-burnt bones, the latter being in rather large quantities on the Continental markets. They contain 45 to 70 per cent of their phosphoric acid soluble in the oxalate. That the phosphoric acid in the two latter named states, of being speedily assimilated in the soil, is known to every farmer. It is not said, however, that they would not profit by being treated with sulphuric acid and converted into bone superphosphate.

Then follow some mineral phosphates, in which 30 to 50 per cent of their phosphoric acid is soluble in the oxalates—as some phosphates of the French Ardennes, in the neighbourhood of Rheims, Dun, Verdun, some phosphates of Cahors, St. Antoine, St. Jean de Laur, Cher, and Lot. All those phosphates are used with excellent effect, only finely ground, for manuring purposes, not only in the neighbourhood of the places named, but even at considerable distances. This proves that their employment was always crowned with decided success; for fresh-broken woodland, turfy ground and soils rich in humic acids, they exercise an excellent effect, even better than superphosphate could do.

In this category is also to be named a phosphate which is found near Mone, in Belgium, in quantity almost inexhaustible. Up to the present time, however, no means have been found of making use of these layers on an extensive scale. This phosphate is found under three forms:—(1.) As phosphatic limestone with 1 to 3 per cent of phosphate of lime. It has been used several years in that region for liming the fields, after being burnt like common limestone. How far the phosphoric acid acts in this manner is very difficult to say. In any case the liming of the fields must have had some success, as it is very generally employed in Belgium for the heavy clay soils. This phosphatic limestone is mostly found at the very surface in layers of 3 to 12 feet in thickness.

(2.) As so-called nodules, a kind of coprolite form in the size of from $\frac{1}{2}$ to 5 inches diameter. These coprolites contain 30 to 40 per cent of phosphate of lime, and are enclosed in a kind of sandy marl containing 5 to 10 per cent of phosphate of lime. The layers have a thickness of 3 to 9 feet, and are for the greater part on the surface. The deeper layers are dug by means of small galleries. The cleaning process of these nodules is carried on in a very simple and rude manner by dry-sifting. They serve, finely ground along with very rich phosphates, for manufacturing superphosphate. A large quantity of them is being sent to England. The chief layers, however, are formed by a kind of hard marl of brownish colour, which is found at a depth of 15 to 20 feet in a thickness of 12 to 20 feet. It contains 20 to 35 per cent phosphate of

lime, and is dug out in about the same manner as the two other kinds named. Of course there are also immense layers of this latter kind which are poorer in phosphate, and up to the present time have not been considered at all.

There are at present two works on the spot to prepare this marl for the market. The one works with a dry setting and sifting process; the other by burning the broken marl pieces in lime-kilns, and washing out a portion of the lime with water. They both make a sandy-looking phosphate of 50 to 55 per cent of phosphate of lime. In either case there remains too much lime and carbonate of lime to enable it to be used alone in the superphosphate manufactory. All these phosphate minerals of Mons contain a mass of Belemnites and other petrified masses of the kind, mostly the marl, which seems to be entirely composed of such. Though the origin of this mineral appears to be of a comparatively recent period, it is for agricultural purposes worthless, with the exception of soils which contain an unusually large quantity of humus and surf. The reason of this is that carbonate of lime, along with the phosphate of lime, enters very largely into the composition of this mineral.

Dr. Petermann, manager of the Belgian Agricultural Stations, has proved by his extremely profound and well-conducted researches* that the above named phosphate of Mons (Ciply) is unprofitable for agricultural purposes, at least under most circumstances. Not only is this phosphate nearly insoluble in different salt solutions, but it is even capable of extricating out of dunghill drainings the phosphoric acid held in solution, and to precipitate it as a tribasic phosphate of lime not soluble in the citrate of ammonia.

From these researches we might conclude that in soils very rich in chalk, as in the Champagne and some parts of England, the plants assimilate the phosphoric acid necessary for their growth as tribasic phosphate of lime. But against this supposition, again, we have the results of Petermann's practical experiments, in which he found that in a poor sandy soil, in which superphosphate was still increasing the harvest very considerably, phosphate of Mons directly employed gave a harvest even inferior to the experiments made without manure phosphate at all. In rich clay soil this astonishing difference was even still more apparent.

The great quantity of carbonate of lime this mineral contains also prevents it from being employed for making superphosphate or precipitated phosphate. A process was proposed to enrich it, which in the laboratory answers well enough if conducted carefully, but on a large scale is impracticable. It is this—to take out the carbonate of lime by means of very weak muriatic acid; but strange to say, there arrives a moment in the process when there is formed precipitated phosphate, while free muriatic acid and carbonate of lime are still present. This precipitate is caused by the carbonate of lime in the solution of phosphate of lime, which latter is dissolved along with the carbonate.

(To be continued.)

New Preparation of Carbolic Acid.—Messrs. F. C. Calvert and Co. have introduced for domestic disinfecting purposes, solidified disinfecting tablets containing 50 per cent of concentrated carbolic acid. The great advantage of this solid form is that it removes the danger of accidental poisoning, and from the high percentage of acid present, the tablets provide, even when dissolved in a considerable quantity of water, a solution capable of arresting all kinds of putrefactive change, and of destroying all the lower forms of insect, vegetable, or germ life.

* "Seconde Note sur les Gisements de Phosphates en Belgique, et particulièrement sur celui de Ciply." Par Dr. A. Petermann, Directeur de la Station Agricole de Gembloux. Brussels, chez Majolez, 1878.

UPON THE DETECTION AND ESTIMATION OF NITROUS ACIDS IN POTABLE WATERS, ACIDS, &c.

By ALBERT R. LEEDS.

I.—WITH META-DIAMIDO-BENZOL.

THIS method depends upon the observation, originally made by P. Griess,* that diamido-benzoic acid is an extremely delicate test for nitrous acid, giving in very dilute solutions of this body an intense yellow colour. Later, Griess proposed to employ, instead of this compound, another related compound, *meta-diamido-benzol*, which is more readily prepared, and, moreover, is twice as sensitive as diamido-benzoic acid. The latter indicates the presence of one-fifth m.grm. N_2O_3 in a litre of water; the former, of one-tenth m.grm., or 1 part in 10 million.† Since *meta-diamido-benzol* is said to be an article of commerce at present there is no need, perhaps, to allude to its preparation. But for those who, like ourselves, were unable to purchase it, it may be of service to know that we were able to prepare it more readily by reduction of the dinitro-benzol by means of tin and hydrochloric acid than by distillation with acetic acid and iron filings.‡ In accordance with the recommendation of Griess, after purifying with animal charcoal, the solution used in testing was acidified with sulphuric acid.

The minutiae essential to employment of the method in quantitative colorimetric determinations have been elaborated by C. Preusse and F. Tiemann,|| who recommend—

- (1.) A solution of *meta-diamido-benzol*, containing 5 grms. per litre. It is acidified with dilute sulphuric acid.
- (2.) Dilute sulphuric acid solution (1:3), for acidifying standard and unknown solutions in process of estimations.
- (3.) A solution of alkaline nitrite, of which 1 c.c. contains 0.01 m.grm. N_2O_3 . 0.0406 gram. pure dry silver nitrite is dissolved in hot water, and decomposed by potassium or sodium chloride. After cooling, the solution is made up to 1 litre, the silver chloride allowed to settle completely, and 100 c.c. of the supernatant liquid diluted to 1 litre.

The rest of the operations are quite similar in character to those made use of in Nesslerising. But instead of using cylinders of colourless glass, as the above-mentioned authors recommend, comparison-tubes and the colour-comparator§ may be more conveniently employed. 100 c.c. of the aqueous solution under examination is introduced into a comparison-tube, 1 c.c. of solution (2) and 1 c.c. of solution (3) added. If the water is not colourless it should first be decolourised by throwing down the colouring matters through formation of an insoluble carbonate of an alkaline earth in the water containing them. If, on agitation with a bulb-stirrer, a caramel-like colour is developed, the trial is to be repeated with 50, 20, 10 c.c. of the solution, previously diluted to 100 c.c. with water (free from nitrous acid). The dilution is sufficient when a satisfactory reaction is attained in the course of one or two minutes. Various amounts of solution (3) are treated in like manner until an identical tint is developed, when the amount of N_2O_3 in the unknown solution becomes, of course, known.

Inasmuch as the tints of colour developed by the *meta-diamido-benzol* are so analogous to those exhibited by a dilute caramel solution that they can be closely copied by the latter, the author uses such a *caramel solution in the glass wedge of the colour-comparator for estimations of nitrous acid*. The same wedge, provided with a different scale, answers for the determination of minute amounts of ammonia by the Nessler reagent.

* Ann. Chem. u. Pharm., cliv., 335.

† Ber. der Deutsch. Chem. Gesell., xi., 623.

‡ "Aromatic Diamines," Hofmann, Proc. Roy. Soc., xi., 518. "Isomeric Diamines," Hofmann, *Ibid.*, xi., 639.

§ Ber. der Deutsch. Chem. Gesell., xi., p. 627.

§ Proc. Amer. Chem. Soc., vol. ii., p. 1.

Amounts of Nitrous Acid found in Potable Waters.

At the time at which these determinations were made, the water of the Passaic, as drawn from the hydrant in the laboratory, contained 0.12 part N_2O_3 in one million. It is of practical importance to compare this result with the amount of nitric acid contained in the same water. A determination of the sum of nitrates and nitrites (calculated as HNO_3) gave, on the same date (May 2, 1879), 2.37 parts in a million. If we regard this amount of nitric acid as an index of previous sewage contamination, its amount is strikingly great as compared with the nitrogenous matters present in the water and still capable of undergoing putrefactive decomposition. The amount of "nitrates and nitrites" contained in the Passaic, May 2, 1879 (determined by reduction with pig-iron), did not differ notably from the amount found Nov. 26, 1877 (by reduction with the copper-zinc couple), when it was 2.22 parts in a million. The ammonia contained in the Passaic water at latter date was 0.02 part; the albuminoid ammonia, 0.215 part in a million. P. Griess has emphatically insisted upon the importance of determining the nitrous acid in potable waters, and their unfitness for household use when they contain nitrous acid.* In cases like the one cited, where the "nitrates and nitrites," so called, exceed the amount of the albuminoid ammonia ten times, there should be no omission to make a separate examination for, and a separate estimation of, the nitrous acid. The rapidity and ease of Nesslerising is such that ammonia and albuminoid ammonia determinations now play the most conspicuous part in reports of water analyses, especially in this country. To accept or condemn from them alone the character of a drinking-water we regard as a practice fraught with error. They should be checked by the results afforded by independent determinations of the nitric and nitrous acids. Fortunately, by aid of improved methods, this can be as readily done as the corresponding estimations of ammonia and albuminoid ammonia, and a skilful water analyst should be able to perform all four within the space of two hours.

We should be very unwilling to say that by reason of the water of the Passaic containing 0.02 part of ammonia and 0.215 part of albuminoid ammonia in a million it was unfit to drink. But these facts, taken in connection with these two others, that it contains at the same time 2.14 parts of nitric acid and 0.12 part of nitrous acid in a million, do, we think, constitute reliable grounds for pronouncing against it. And we regard the two facts last mentioned of more weight in making up this opinion than the two first. Water flowing from a granitic water-shed and over alluvial country, as the Passaic River does, could not readily obtain this percentage of nitrates from inorganic sources—it has most probably come from the oxidation of nitrogenous animal and vegetable matters. And this opinion is confirmed by finding an amount of nitrous acid which is *one-half the total amount of the ammonia-yielding bodies present*, and represents how large a portion of these last are still in a state of transition from their evanescent putrescible form, as albuminoid ammonia, to their permanent stage, as nitric acid.

The water of the Passaic has been taken as an illustration because it happens to be the only potable water which at the present time I have occasion to analyse. But having alluded to it, I trust the importance of the subject will excuse my saying here that the enforced use of water unquestionably so contaminated as this is a crime committed against the health and happiness of a quarter of a million of people.—*Journal of the American Chemical Society.*

"Malco."—This nostrum, vaunted as a prophylactic against the plague and all infectious diseases, consists of 2.5 grms. ammonium carbonate, scented with oil of roses and carbolic acid. It is inclosed in a small leather case, and is worn suspended round the neck!—*Pharm. Zeitung.*

* *Ann. Chem. Pharm.*, cliv., 336.

ANALYSES OF THE
WATERS OF LAKE THIRLMERE AND THE
RIVER VYRNYW, THE NEW SOURCES
OF WATER-SUPPLY FOR
MANCHESTER AND FOR LIVERPOOL.

By C. ESTCOURT, F.C.S.

IN December, 1877, I visited Lake Thirlmere and took samples of the water, one at the head of the lake, and one at its present outlet. I had intended visiting the lake during the ensuing summer for the purpose of observing the difference between the two seasons, but circumstances prevented me from carrying out this intention.

I find that since then Messrs. Grimshaw have analysed samples taken about the end of the summer of last year (namely, August, 1878), see *CHEMICAL NEWS*, vol. xxxviii., p. 216, and it will probably not be uninteresting to compare the results of the analyses of the winter and summer samples with each other.

During the present summer I visited the River Vyrnyw, at Llanywdden, in Montgomeryshire, at which place the river is for four miles to be converted into a lake for the purpose of supply-water to Liverpool; I procured a sample of the water at a spot within the area of the intended lake. The following are the results of my analyses together with those of Messrs. Grimshaw, of Thirlmere.

Per Gallon.	Thirlmere Water Unfiltered.					
	Estcourt. Head of Lake.	Grimshaw. Upper.	Estcourt. Outlet.	Grimshaw. Lower.	Vyrnyw Filtered.	Estcourt.
Total solid matter ..	2.25	2.20	3.10	3.15	5.16	
Mineral matter ..	1.10	1.50	1.15	1.40	1.46	
Loss on ignition ..	1.15	0.70	1.95	1.75	3.70	
Hardness, all permanent	2.25	1.00	1.50	1.00	2.75	
Chlorine	0.58	0.42	0.44	0.70	0.41	
Ureal ammonia ..			0.0009	0.0021	0.0154	
Albuminoid ammonia			0.0042	0.0042	0.0056	
Colour	none	none	none	none	Pale yellow.	

It will be perceived that all analyses of Thirlmere water gave results within the limits of extreme purity. The difference in the degree of hardness, shown in the analyses of Messrs. Grimshaw and myself, may arise from the use of a different standard solution. One c.c. of the soap solution used in my laboratory is required to produce a lather when 70 c.c. distilled water are shaken with it. Thus the hardness in my samples due to mineral matter will be 1.25° and 0.50° respectively.

In the Vyrnyw water the amount of organic matter which it contains is indicated readily by its peaty colour, and it will be observed the ureal ammonia is high. The unfiltered water contained brownish floating particles which could not be got rid of by decantation, therefore filtration was necessary. After the filtered water had been standing for a fortnight the colour had not diminished in depth.

ON A METHOD FOR THE DETERMINATION
OF PHOSPHORIC ACID.*

By S. W. JOHNSON and E. H. JENKINS.

OTTO long ago proposed a method for the separation of phosphoric acid from iron and aluminum which was based on the fact that ammonium tartrate will prevent the precipitation of the hydrates and phosphates of the metals in alkaline solutions, but does not prevent the precipitation of ammonia-magnesium phosphate.

* From the Report of the Connecticut Agricultural Experiment Station for 1878.

W. Mayer* has shown that unless a large amount of ammonium salts is present in solution, basic magnesium tartrate will also be precipitated with the phosphate. He prepared a solution for use in the determination of phosphoric acid in which the amounts of magnesium and ammonium salts were in such relation that a precipitation of basic magnesium tartrate was not to be feared. Such a solution cannot be used, however, for the direct gravimetric determination of phosphoric acid in the presence of much lime, because neutral calcium tartrate is apt to be precipitated with the ammonio-magnesium phosphate.

F. Stolbat† has shown that pure ammonio-magnesium phosphate can be determined by titration as well as by weighing, one molecule of the pure salt requiring two molecules of hydrochloric acid to destroy its alkaline reaction. Advantage has been taken of these observations in devising a plan of operating which should meet the want felt for a rapid and accurate method of determining phosphoric acid in commercial fertilisers. The standard acid used in other volumetric work answers perfectly for this. A strong, nearly saturated, solution of ammonium tartrate, free from carbonic acid, and a solution of some magnesium salt, are also necessary. The latter is prepared by dissolving 70 grms. of magnesium sulphate and 195 grms. ammonium chloride in 1 litre of water. 10 c.c. of this solution contain twice the amount of magnesium necessary to precipitate 0.1 gm. phosphoric acid (P_2O_5). A suitable amount of the phosphate (in most cases 1 gm. is a convenient quantity) is dissolved in hydrochloric acid, the solution nearly neutralised with ammonia, and ammonium tartrate solution is added, 10 c.c. at a time, till the solution remains perfectly clear when slightly alkaline. Add a suitable quantity of the magnesium mixture, and either stir vigorously with a rod, or, if the precipitation is made in an assay-flask, as it can be very conveniently, shake occasionally. When the precipitation is nearly complete add enough ammonia to make it very strongly alkaline, and let it stand six to twelve hours. It can then be filtered, preferably on the pump, and washed with equal parts of strong alcohol, 85 to 90 per cent, and water. No pains are taken to detach the precipitate from the glass. When the dish and precipitate are washed until the washings no longer react alkaline, the filter and precipitate are brought back into the beaker or flask, a little water and a few drops of cochineal tincture are added, and it is titrated. This is best done by adding an excess of standard acid at once, stirring so that all the precipitate shall be wetted with it, and after it has stood a few minutes, measuring back with standard alkali.

The results given below (mostly duplicated) indicate the degree of accuracy to be expected.

	Determined by use of Ammonium Molybdate.	Determined by the Method just described.
Superphosphates, soluble } phosphoric acid	8.92 to 8.96	8.83 to 8.91
Ditto, ditto.. ..	8.31	8.32
Ditto, ditto.. ..	11.89 ,, 11.95	11.83 ,, 11.95
Ditto, ditto.. ..	5.14 ,, 5.08	5.07 ,, 5.11
Ditto, ditto.. ..	6.78	6.68 ,, 6.84
Ditto, ditto.. ..	5.63 ,, 5.65	5.61 ,, 5.63
Superphosphates, total } phosphoric acid	9.21 ,, 9.28	9.22 ,, 9.32
Ditto, ditto.. ..	10.70 ,, 10.72	10.76 ,, 10.87
†Ditto, ditto.. ..	16.66 ,, 16.66	16.55 ,, 16.65
‡Ditto, ditto.. ..	13.94	13.90 ,, 14.05
Hair manure, ditto ..	2.23	2.19
Bone ditto, ditto ..	21.90 ,, 21.90	21.87 ,, 21.75
Ditto, ditto.. ..	13.21 ,, 13.40	13.19 ,, 13.33
Ditto, ditto.. ..	22.57 ,, 22.62	22.57 ,, 22.76
Ditto, ditto.. ..	21.90 ,, 21.90	21.87 ,, 21.75
Fish scrap ditto, ditto ..	6.17 ,, 6.27	6.17 ,, 6.28

* Ann. Chem. Pharm. 101, 164.

† Zeitschr. Chemie, xvi., 100.

‡ Navassa superphosphates containing soluble iron and aluminum phosphates.

Tricalcic phosphate with 6.44 p.c. water	Calculated.	42.85	42.79
The above (0.5 gm.) with 0.22 gm. iron			
in form of ferric chloride		42.85	42.79

In the case of a few phosphatic materials rich in phosphoric acid, larger discrepancies than any above given have been occasionally encountered, and it is proposed to give them further attention, although it is our impression that they were accidental and do not invalidate the accuracy of the method.

This process requires less than half the time and labour necessary for the molybdic method, is scarcely less accurate, and appears to be generally applicable.

Some investigations, not completed as yet, lead us to hope that ammonium tartrate may be successfully substituted for ammonium citrate for bringing precipitated or reverted phosphates into solution. This step would still further simplify the analyses of superphosphates, since the entire phosphoric acid, soluble, reverted, and insoluble, could be quickly estimated in a single portion.

Our investigations have also demonstrated that while ammonio-magnesium phosphate is totally insoluble in a large excess of ammonium tartrate, it is soluble in excess of ammonium citrate, and hence all methods based on the use of citric acid are faulty.

It is a fact also that ammonio-magnesium phosphate is largely soluble in ferric and aluminic solutions, containing *insufficient* ammonium tartrate.

It is therefore necessary in presence of iron to add ammonium tartrate more than enough to produce a reddish yellow solution, enough, in fact, to make a greenish yellow solution, as Otto has indicated. A similar excess of ammonium tartrate is also requisite in presence of aluminium, and while there is no colour-indication of the suitable quantity, a large excess does not appear to retain ammonio-magnesium phosphate in solution, unless the liquids are too concentrated.—*American Chemical Journal*.

NOTE ON THE DETERMINATION OF SILICON IN PIG-IRON AND STEEL.*

By THOMAS M. DROWN.

In experimenting in connection with Mr. P. W. Shimer (now chemist of the Thomas Iron Company, Hokenau, Pa.) on methods for the determination of silicon in pig-iron, in order to find one which should be accurate and yet give results in a few hours, I have adopted the following procedure, which, as far as my experience goes, leaves nothing to be desired.

About 1 gm. of pig-iron or steel is treated in a platinum or porcelain dish with 25 c.c. of nitric acid (sp. gr. 1.2). When action has ceased 25 to 30 c.c. of dilute sulphuric acid (one of acid and three of water) are added, and heat applied until the nitric acid is nearly or quite driven off. The heat of a water-bath is sufficient, though the process may be hastened by heating higher on a sand-bath. Water is then cautiously added (as soon as the free sulphuric acid is sufficiently cool) and the contents of the dish heated until the crystals of ferric sulphate are completely dissolved. The solution is then filtered *as hot as possible*, the residue washed first with hot water then with 25 to 30 c.c. of hydrochloric acid (sp. gr. 1.12), and finally with hot water. After drying and igniting the silica will be found to be snow-white and granular.

The following are some results obtained by this method compared, in some instances, with the older method of treatment with nitric acid, evaporation to dryness, heating to 150° C. for several hours, dissolving out the iron in hydrochloric acid, and filtering off from the insoluble residue, which is dried and ignited, and the resulting impure silica fused with alkaline carbonates.

The letters denote different samples of pig-iron.

* Read before the American Institute of Mining Engineers.

Per cent of Silicon.							
	A.	B.	C.	D.	E.	F.	G. Bessemer Steel.
Old method—	2.64	2.46		1.45	1.65		0.672
New method—							
1.	2.70	2.47	1.13	1.63	1.53†	1.66	2.30 0.676
2.	2.68	2.47	1.18	1.62	1.51†	1.68	2.30 0.672
3.	2.81	2.47		1.65	1.51†	1.72	2.50 0.672
4.					1.63‡	1.70	2.47
5.					1.65‡		2.46
6.					1.65‡		

Some incidental results obtained in developing this process have enough interest to be worthy of record. Treatment of pig-iron with concentrated sulphuric acid, heating till fumes arise, diluting with water, and filtering after all action has ceased, gives a silica which is seldom pure, and yet the results are considerably too low.

Treatment with dilute sulphuric acid and evaporation till the acid fumes in the air, then filtering after dilution, gives occasionally results which are accurate; but this method is uncertain, depending on the fineness of the borings and character and composition of the pig-iron. The silica obtained is seldom white. The following are some results obtained in this way:—

Per cent of Silicon.				
	A.	B.	C.	D.
Old method ..	1.02	1.64	2.64	3.85
New method, 1..	1.05	1.73	3.00	3.88
" " 2..	1.05	1.69	2.98	3.91
" " 3..	1.05	1.70	2.97	
" " 4..			3.01	

Treatment in platinum dishes gave very slightly lower results than porcelain dishes.

If, after treatment with dilute sulphuric acid, the solution is filtered off from the residue without concentration of the acid, it is found that about one-half of the silicon is in the solution and the other half in the residue; when nitric acid is used and the solution filtered off as soon as all action has ceased, it is found that about two-thirds of the silicon is in the solution and one-third in the residue; and with hydrochloric acid, about one-third goes into the solution and two-thirds remain in the residue. It is not probable that there is any precise ratio existing between the amount of silicon dissolved and the amount in the residue in the case of any one of the acids, the ratio being doubtless variable and depending on the concentration of the acid, the time of action, and the temperature; yet the marked difference in the action of the three acids in this respect is interesting.

The washing with hydrochloric acid of the residue obtained by the action of nitric and sulphuric acids on pig-iron is in most cases necessary. Thus there was obtained from the pig-iron when water only was used for washing, 2.67 per cent of silicon against 2.52 when washed with hydrochloric acid; and in another sample, 2.10 per cent against 1.70 per cent.

Although the results obtained with hydrochloric acid for the original solution of the iron show, as far as the experiments go, as good results as those obtained with nitric acid, yet I prefer the nitric acid treatment on account of the silica obtained being compact and granular, while the use of hydrochloric acid, and also of sulphuric acid alone, yields a silica which is light and flaky.—*American Chemical Journal*.

According to the *Pharm. Zeitung* American honey can be distinguished from that of European origin by the different character of the pollen grains.

* Not washed with hydrochloric acid.

† In these analyses hydrochloric acid was used after the addition of nitric acid, and was not completely driven off.

‡ Hydrochloric acid was used for solution instead of nitric acid.

NOTICES OF BOOKS.

Copyright and Patents for Inventions. Vol. I., Copyright. By R. A. MACFIE. Edinburgh: T. and T. Clark.

By "reform" of existing patent legislation we have always understood increased facilities and decreased burdens for inventors, so that England may be no longer, as at present, heavily handicapped in its industrial competition with America. Here, however, we find the same "reformers of the actual legislation with respect to books and inventions" coolly allotted to men who are labouring in the very opposite direction, and of whom we are told that "but a few aim at abolishing privilege (!) indiscriminately and entirely."

The author persists in ranking copyright and patent-right among monopolies, and remarks that "they have been spared or instituted when other kinds of exclusive privileges were condemned and abandoned by all statesmen." Might not this very fact suggest to Mr. Macfie and his fellow-believers that these "privileges" have, in reality, no community of nature with monopolies, and were merely confounded with such by the ignorance of our forefathers, just as they confounded whales with fishes and bats with birds? The monopolist is a man who takes something from the public and appropriates it exclusively to himself. The inventor does nothing at all analogous; he makes an exclusive claim to something which, but for him, would have been non-existent. In any case the public, therefore, would be no loser. That if he had not come upon the novelty someone else might is beside the question, for then that other person would have been the inventor. Mr. Macfie's sole ground for denying to a novel invention or a new book the title to rank as property is because it is less easily defined than land, and is not "visible and palpable"—a somewhat flimsy argument. If the author would look round he might meet with things not a few neither visible nor tangible, and certainly not the creation of their holders, which the law recognises as property. Thus if a number of persons associate themselves together, and propose to supply any town with gas or water, the Legislature will give them a perpetual property, not merely in their works, plant, mains, and other "visible and palpable" things, but in the sole right to supply such town. We even suspect that if it were proposed to supersede gas by, say, the electric light, a cry for "compensation" would be raised on behalf of the gas companies. It may, however, scarcely seem fair to discuss the question of patents till the second volume of the work has appeared, though, as far as we can judge from the preface to the volume before us, the author has little to present, save fallacies, which are now very far worn. We will merely remind him that without patents there will be no inventions, and without inventions no trade.

As regards literary property, it seems to be attacked chiefly on the plea that it "enhances their price beyond all bounds." No doubt if the author could be compelled to work gratuitously books would be somewhat cheaper. But why is he alone to be subject to such demands? If the paper-maker, the printer, the bookbinder, and the publisher would all forego wages and profits a still further reduction could be made. Perhaps some of these parties to the transaction could afford to dispense with their emoluments quite as well as the author could forego his "copy-money." If we bear in mind that a scientific book is often the outcome of years of study and research, we shall find that this "copy-money" would often not amount to a stone-breaker's wages for the time consumed.*

The high prices of scientific books in England, in comparison with those prevailing in some other countries seem to have their origin in the following circumstances. Book buyers may be divided into three classes, which we will distinguish as A, B, and C. Class A is composed

* Not unfrequently an author receives nothing at all.

of men who care little for the contents of a book, but who will buy any work that "makes a noise," provided it is showily got up. Class B consists of men able to appreciate such books, and who would purchase them in any case. Class C is composed of poor students and men of science in humble circumstances who cannot afford typographical luxuries, though they would be purchasers of a cheaply got-up work. In England, $a + b$ are together much more numerous than $b + c$. Hence the publisher finds it to his interest to bring out every new work of interest in an expensive style. In Germany the Class C is so numerous that scientific works are published in a cheap form for their use. The scheme of publishing on royalty is open to the difficulty that it might not suit the trade. Except an author has a very high name, he finds it no easy matter to arrange with a publisher, and the latter would in most cases decline the transaction altogether unless the copyright were, as now, handed over to him exclusively.

We regret to see a man whose intentions are good labouring perseveringly and ably in opposition to what we consider to be the best interests of his country.

Is the Atomic Weight of Antimony Sb, 120, or 122?
A Reply by Dr. F. KESSLER. Bochum: Ad. Stumpf.

THE author, after referring to the atomic weight of antimony as given by Berzelius, $Sb = 120$, and now universally recognised as too high, cites the results obtained by five chemists who have investigated this question. Schneider gives the value as 120.30, Dexter 122.33, Dumas 122.00, the author himself in his earlier researches 122.37, and Cooke 120.00. The material selected by the first-named authority, native antimony-glance, was very probably impure. The results obtained by Cooke differ widely among themselves, and it seems difficult to say why he finally gives the preference to 120. Dexter's result, 122.03, is the mean of 11 determinations, the highest of which showed 122.18 and the lowest 121.93. Hence we see that three chemists, proceeding by totally distinct methods, obtained numbers which arrange themselves very closely round the whole number 122, *i.e.* :—

Dexter	..	..	..	..	122.03
Dumas	..	..	..	..	121.95
The Author	..	..	..	..	121.90

Dr. Kessler, whilst accepting 122 as the nearest approximation to the true atomic weight of antimony, has no intention, however, of declaring himself a supporter of Prout's hypothesis.

Proceedings of the Aberdeenshire Agricultural Association. Report of the Third Annual General Meeting, Season 1878.

THE Aberdeenshire Agricultural Association, which is presided over by the Marquis of Huntly, although only three years old, appears to have already largely contributed to our knowledge of the scientific principles of agriculture. Mr. T. Jamieson, the chemist of the Association, has evidently concentrated the whole of his energies upon turnips, which appear to be one of the staple agricultural productions of the ungrateful soil of Aberdeenshire. We wish we had sufficient space to give a summary of Mr. Jamieson's very able report, which in clearness and conciseness is a model of what such a report should be; we can only give the conclusions which he has arrived at after three years' experiments. They are as follows:—First, that calcic phosphates, whether of animal or mineral origin, decidedly increase the turnip crop; secondly, that soluble phosphate is not superior to insoluble phosphate to the extent that is generally supposed; thirdly, that nitrogenous manures when used alone have little effect on the crop, but when used in conjunction with insoluble phosphates they in-

crease it; fourthly, that the addition of nitrogen to soluble phosphates does not increase the dry matter of the crop; fifthly, that there is no difference between the effects of equal quantities of nitrogen in sodic nitrate and ammoniac sulphate; and, lastly, that fineness of division seems nearly as effective in increasing the crop as the addition of nitrogenous manures. Hence the most economical phosphatic manure for turnips is probably insoluble calcic phosphate from any source, animal or mineral, ground down to an impalpable powder. The tables annexed to the report contain full details of the methods adopted, the analyses of the soil and manures employed, the size of the plots, the number of turnips, the weight of their leaves and bulbs, specific gravity, and percentage of solids in the bulbs. Mr. Jamieson and his able assistants fully deserve all the praise bestowed on them by Mr. J. W. Barclay, M.P., for the carefulness and completeness with which the experiments have been carried out.

We should be only too glad to see every county in England, Ireland, and Scotland follow the excellent example set by hard-headed, hard-soiled Aberdeenshire.

The Retrospect of Medicine, &c. Edited by W. BRAITHWAITE, M.D., and JAMES BRAITHWAITE. Vol. lxxix., January to June, 1879. London: Simpkin and Marshall.

THE present issue of this valuable compilation is as usual almost devoid of interest to the chemist, and shows once more conclusively the persistent manner in which the members of the medical profession neglect to search the rich armoury of yet unsheathed weapons placed at their service by chemical science. We do not, however, blame medical men, but the wretched cram system which obliges them to painfully coach up a subject which when studied in the laboratory is delightful, but when learned from books must, we imagine, be especially distasteful. Hence the medical man who is also a chemist is a *rara avis* indeed. One of the only novelties likely to interest our readers is the introduction of pyrogallic acid into medicine by Dr. Vesey, of Dublin, who administers it in grain doses in hæmoptysis and other cases of internal hæmorrhage. The dose is small, and it does not derange the stomach like its congeners, gallic and tannic acids, nor does it cause vomiting as iron and ergot often do. We should imagine that the best way of exhibiting it would be in the form of a spray. The use of arsenic as a cardiac tonic, by Dr. Steward Lockie, of citrate of caffeine as a diuretic, and of sodium ethylate as a topical caustic, may also be mentioned among matters of chemical importance. There is an interesting article on the action of anæsthetics, by Drs. Coats, Ramsay, and McKendrick; and on dichloride of ethylene as an anæsthetic, by the last named investigator, which are models of what such researches should be.

CORRESPONDENCE.

JOHNSTONE'S BAROMETER PUMP.

To the Editor of the Chemical News.

SIR,—In the *CHEMICAL NEWS*, vol. xl., p. 22, under the heading of "Notices of Books," there appears a review upon C. Greville Williams's "Supplement to a Handbook of Chemical Manipulation," in which the reviewer draws attention to what he calls an excellent account of what is at the present time called the Sprengel pump.

Circumstances have, until now, prevented my calling attention to an erroneous idea which is in circulation at the present moment, viz., that the invention of what is called the Bunsen and Sprengel pumps, "*solely and only belong to Dr. Sprengel.*" (See *CHEMICAL NEWS*, vol. xxvii.,

p. 49). The above statement made by Prof. Bunsen, and also the statement by Mr. Williams, that the Bunsen pump, "is an outcome from Sprengel's discovery," I give a decided denial to, as neither Dr. Sprengel, or Prof. Bunsen has any claim whatever to the discovery. Dr. Sprengel, in his original paper read before the Chemical Society, made the following statement:—"The main fact which I have established in this paper may be shortly stated to be that, if a liquid be allowed to run down a tube, to the upper part of which a receiver is attached by means of a lateral tube, and if the height at which the receiver is attached be not less than that of the column of the liquid which can be supported by the atmospheric pressure, a vacuum will be formed in the receiver, minus the tension of the liquid employed. To the originality of the above statement Dr. Sprengel has no claim whatever, and I now give my reasons for so stating. In the year 1845 James Johnstone (my father) had a patent granted to him under the following title "Specification of the Patent granted to James Johnstone, of Willow Park, Greenock, Engineer, for New and Improved Process and Machinery for Making and Refining Sugar.—Sealed January 31st, 1845." A full description of the patent will be found in "The Repertory of Patent Inventions for 1845" and I extract the following paragraph from the specification of the patent, in which Mr. J. Johnstone says—"I attach a pipe to the bottom of a vacuum pan, so that the saccharine fluid within the pan can freely enter the pipe, and the pan may be emptied thereby. I call this the barometer pipe; it must be made of such length that the weight of the saccharine fluid within it shall counterbalance or rather exceed the weight of the atmosphere, so that when a stopcock at the bottom of the pipe is opened the saccharine fluid within the pan will, of its own weight, rush out by the pipe till the pan is emptied; but the pipe will remain filled with the saccharine fluid in the same manner as a common barometer tube remains full of mercury. The pipe for this purpose will be upwards of twenty-five feet long; the length of the pipe required depends on the specific gravity of the saccharine fluids after they are boiled; the heavier the fluids the shorter the pipe may be made. The preceding is the least expensive method of emptying a vacuum pan without destroying the vacuum, but as it may not always be convenient, instead of the barometer pipe, a pump may be attached to the pan, for the purpose of removing the saccharine fluids without destroying the vacuum; but I prefer the barometer pipe when it can be conveniently used. I claim the making and using of vacuum pans with apparatus attached, so that the contents of the pans may be withdrawn without the vacuum within the pans being destroyed." I think I have now proved conclusively that those people who persist in speaking of the Johnstone barometer pump as the Sprengel or Bunsen pump have now no warrant for doing so, as the invention belongs solely and only to James Johnstone, he having used mercury as well as saccharine fluids for creating vacuum.—I am, &c.,

WILLIAM JOHNSTONE.

Lowther Hill, Forest Hill, July 15, 1879.

[The quotations which Mr. Johnstone gives from his father's specification show that the "Johnstone barometer pump" is an entirely different instrument from the "Sprengel pump."—Ed. C. N.]

Alloys of Lead and Antimony.—J. de Jussieu.—These alloys melt at 355°, and only give off vapours at a red heat. They dissolve in melted lead and crystallise by liqutation in the rhombohedral system. They are capable of being tempered, the effect of which operation is very superficial. They are unstable and are readily decomposed by the action of heat, lead being set at liberty, and a compound richer in antimony being formed.—*Les Mondes*.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Justus Liebig's Annalen der Chemie,
Band 196, Heft 3.

Certain Sulphur Oxychlorides.—The author, referring to certain experiments by Rose on this subject, considers that the "mother-liquor" of this chemist consisted of the decomposition-products of a small quantity of $\text{S}_2\text{O}_3\text{Cl}_4$ which was formed, and that everything else was unchanged sulphuric anhydride. He has also examined a number of complex selenium compounds, such as sulpho-selen-oxy-tetra-chloride, and the behaviour of sulphuryl-hydroxyl-chloride with the chlorides of titanium, antimony, tin, and silicon.

Colouring Matters and the Glycoside-sugar of Persian Berries.—C. Liebermann and O. Hörmann.—The authors do not consider the colouring matters of quercitron, berries, and fustic as identical. Berries, especially those from *Rhamnus infectorius* and *tinctoria*, even when kept for many years, still contain about 12 per cent pigment glucosides in addition to a little free colouring matter. The glucoside less freely soluble in alcohol is Kane's xanthorhamnin. The more soluble, Schützenberger's β -rhamnegin cannot be regarded as fully established. Xanthorhamnin is split up by acids into rhamnetin and isodulcite, the sugar formed on splitting up quercitrin. Rhamnetin must have Schützenberger's formula, $\text{C}_{12}\text{H}_8\text{O}_3(\text{OH})_2$. Xanthorhamnin and rhamnetin are not identical with quercitrin and quercetin. Lefort's rhamnin is formed by the action of a ferment upon one of the glucosides, and is still a glucoside itself.

On Azophenols.—P. Weselsky and R. Benedikt.—An examination of para-, meta-, and ortho-azophenol and certain of their derivatives.

Action of Water upon the Haloid Compounds of the Alcohol Radicles.—Gustav Niederist.—The author's researches confirm his former opinion that the haloid compounds of the univalent and bivalent radicles, on treatment with excess of water, can almost always be converted into alcohols.

Band 197, Heft 1.

Boiling-points of the Esters and Ether-esters of Oxy-acids.—Dr. L. Schreiner.—Every oxy-acid esters boils almost at the same temperature as its ethyl-ether. If an oxy-acid ester is converted into the corresponding methyl-ether, the boiling-point sinks as when a normal alcohol is converted into methyl-ether.

Methyl- and Dimethyl-diacetonamin.—Dr. Th. Götschmann.—From the facts arrived at it appears that organic amin bases also can act upon aceton with a formation of keton bases, the conversion ensuing in the same manner as with ammonia.

Contribution to the Knowledge of Quinamin.—A. C. Oudemans, jr.—The author gives the preference to the formula first proposed for this compound by Hesse, $\text{C}_{20}\text{H}_{26}\text{N}_2\text{O}_2$. He describes the following reactions as characteristic, even for very small quantities. If a drop of the solution of a salt of quinamin is cautiously let flow upon concentrated sulphuric acid containing a trace of nitric acid, there appears, at the point where the two liquids come in contact, a chestnut-brown colour if the liquid is concentrated, but if dilute a rich orange. If the whole mixture is then gradually diluted with water it becomes purple and ultimately a faint rose-red. If a few letters are written upon common thick white paper with a goose-quill dipped in a moderately concentrated solution of quinamin in a slight excess of sulphuric acid, and if

the paper is laid with the writing downwards over a watch-glass containing a little strong sulphuric acid and a few granules of potassium chlorate, the characters in a few seconds turn brown or olive. If the paper is removed the letters ultimately turn to a rose colour. If a considerable excess of strong sulphuric acid is added to the quipamin solution prior to the action of chloric peroxide, the letters turn olive more rapidly, and on exposure to the air become sky-blue or deep blue according to concentration. The reaction with chloric peroxide does not occur with quinin, quinidin, cinchonin, cinchonidin, quinicin, and cinchonidin.

Communications from the Laboratory of the University of Kasan.—These consist of a memoir by C. Rjabinin and A. Saytzeff on Diallyl-iso-propyl-carbinol, and a paper by A. Semljanitzin and A. Saytzeff on Oxvalerianic Acid obtained from Allyl-dimethyl-carbinol.

Behaviour of Certain Nitro Compounds with Sulphuretted Hydrogen.—F. Beilstein and A. Kurbatow. —Tri-substituted chlor-nitro-benzols of unsymmetrical structure are not reduced by sulphuretted hydrogen.

Researches on Compounds of the Camphor Group (Sixth Dissertation).—J. Kachler.—An examination of Borneo-camphor, its components, and behaviour with reagents.

On Glycyrrhizin.—J. Habermann.—In the liquorice root there exists a peculiar nitrogenous acid in a saline state. The acid is tribasic, and yields both acid and neutral salts, among which the acid potassium and ammonium salts are especially distinguished by their tendency to crystallise and by their sweetness.

A Camphen Derived from Camphor and the Synthesis of its Homologues.—F. V. Spitzer.—Not adapted for useful abstraction.

Band 197, Heft 2, 1879.

Experimental Researches on Hydrogen Peroxide.—E. Schöne.—Sixth Treatise:—Behaviour of Hydrogen Peroxide with the Galvanic Current.—The author infers from his experiments that hydrogen peroxide is not an electrolyte, but that in its acidulated solutions the water (or more accurately the acid) alone is subject to electrolysis, the decomposition of the peroxide in the electrolytic liquid being a secondary process, consisting herein that both of the electrolytic products of the decomposition of the water in the nascent state exert a reducing action upon the hydrogen peroxide present around the electrodes. A formation of hydrogen peroxide does not take place during the electrolysis of water. The author points out that permanganates, chromic acid, and lime-water as reagents for hydrogen peroxide require to be applied with caution.

The Two Isomeric Bromides, $C_3H_5Br_2$.—E. Erlenmeyer.—The most favourable conditions for the formation of trimethylen bromide are the maintenance of the largest possible proportion of dry hydrobromic acid to the dry allyl-bromide, till the reaction is completed; secondly, a temperature of 30° to 40° .

Oxychlorides and Chlorides of Tungsten.—Hugo Schiff.—A comment on Teclu's memoir (*Annalen*, Bd. 187, p. 255).

Influence of Isomerism of Alcohols and Acids upon the Formation of Compound Ethers.—N. Menshutkin.—This memoir is not capable of useful abstraction.

On Cinchonidin.—Zd. H. Skraup and C. Vortmann.—An account of the platinum double salt; of cinchonidin hydrochlorate and the neutral sulphate; of the oxidation of cinchonidin with potassium permanganate; and of the preparation and properties of cinchotenidin.

Action of Oxidising Agents upon the Hydrocarbons of the Series C_nH_{2n} .—Dr. O. and F. Zeidler.—The action of the various oxidising agents differs quantitatively, not qualitatively. Potassium permanganate,

especially in alkaline solutions, acts more energetically and gives a better yield of bibasic acids, whilst chromic acid and potassium bichromate along with sulphuric acid give chiefly monobasic acids along with small quantities of bibasic acids. The acids formed are, however, in both cases the same.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 4, 1879.

Preparation of Divaleryl.—J. W. Brühl.—The author has succeeded in obtaining divaleryl by a process analogous to that of Freund for the production of dibutyl.

Succinyl Compounds of the Toluydina.—G. v. Bechi.—The author has obtained certain derivatives of ortho-toluy-succinimid and the corresponding compounds of the para series.

Action of Dehydrating Bodies upon Camphoric Acid and its Amides.—M. Ballo.—The bulk of the camphoric acid when acted on by dehydrating bodies is decomposed, as is oxalic acid under the same circumstances. A small portion undergoes a more complicated decomposition with the formation of camphoterpen, $C_{20}H_{32}$.

Hydrocarbons obtained by the Action of Chloromethyl upon Benzol in Presence of Aluminium Chloride.—E. Ador and A. Rilliet.—A description of tri-, tetra-, penta-, and hexa-methyl-benzol.

The Photographic Observation of the Oxygen Spectrum.—Hermann W. Vogel.—The spectrum showed the lines described by Paalzow, O_γ between b and E , O_δ close by F , and O_ϵ between F and G . A band of great intensity, which the author names O_η , lies near h . It is sharply defined towards the red end of the spectrum, but shades away towards the violet extremity. A double band, O_θ , near G , has the same character. In the spectrum of hydrogen the three already-known hydrogen lines in the blue and the violet were seen very distinctly, and also the red line, H_α , coincident with C of the sun. The fourth hydrogen line, coinciding with " h " of the sun, was observed with the naked eye by Paalzow and the author, on the application of the simple induction-current, in opposition to the assertion of Lockyer that it is only visible at very high temperatures. Upon this assertion he founds in part his supposition of the decomposition of hydrogen at elevated temperatures.

Oxidation of Lævulic Acid.—B. Tollens.—The products of oxidation are the succinic, acetic, oxalic, formic (?), carbonic, and hydrocyanic acids.

On Phenyl-phosphin.—A. Michaelis and E. Dittler.—The authors describe experiments on the preparation from phenyl-phosphin of carbonyl-phosphin and carbyl-phosphin.

A New Carbazol from Coal-tar.—C. Graebe and W. Knecht.—This compound, for which the authors propose no name, is represented by the formula $C_{12}H_{11}N$. It melts at 330° , dissolves scarcely if at all in cold alcohol and benzol, sparingly in boiling benzol and glacial acetic acid. It is moderately soluble in hot aniline. The solution has an intense blue fluorescence.

Certain Methyl-esters of the Propionic and Butyric Acid Groups.—G. W. A. Kahlbaum.—A table of boiling-points, specific gravities, and indices of refraction.

Behaviour of Dibrom-succinic Acid with Water.—E. Bandrowski.—This acid and its isomer are decomposed on boiling with water.

Correction in a Communication on Atmospheric Hydrogen Peroxide.—E. Schöne.—The figures in the last column of "Table A" express the hydrogen peroxide in 1000 cubic metres of air and not in 1000 c.c.

Action of Chlorine Gas on the Hydrates of Baryta and Strontia.—J. Konig-Weisberg.—Barium hydroude takes up no chlorine. The absorption of chlorine depends solely upon an excess of water.

Synthesis of Isatin.—L. Claisen and J. Shadwell.—Cyanide of ortho-nitro-benzoic acid is transformed into a ketonic acid, and the latter again into the amide compound by limited reduction. The next stages are ortho-nitro-phenyl-glyoxylate of potassium, isatate of potassium and isatin.

Decomposition of Tyrosin by Putrefaction.—Th. Weyl.—The author has investigated the production of phenol during the putrefaction of tyrosin. In one case he appears to have obtained para-cresol.

Determination of the Vapour-densities of the Three Isomeric Dinaphthyls.—Watson Smith.—These are $\alpha\alpha=8.70$; $\beta\beta=8.73$; $\alpha\beta=8.77$.

Existence of Nitrous Anhydride as Vapour.—G. Lunge.—Nitrous anhydride partially dissociated by evaporation, but a complete dissociation can be reached neither by mixture with a great excess of air nor by higher temperatures. In most cases a considerable residue, often as much as three-fourths, remains undecomposed, and must exist as nitrous anhydride in the form of vapour. An increasing tendency to dissociation with increased excess of air cannot be denied. The effect of temperature is not demonstrated. Nitrogen trioxide can subsist as vapour even at 150° , but it has in this state a tendency to dissociation, which is increased by the presence of air.

Bulletin de la Société Chimique de Paris,
No. 11, June 5, 1879.

Action of Ammoniacal Salts upon certain Metallic Sulphides and Application of the Facts Observed in Analysis.—Ph. de Clermont.—Already noticed.

Observations on Sulphurous Baths.—Ph. de Clermont and J. Frommel.—The authors believe that the action of certain mineral baths may be explained by means of the electric currents which take place in them.

On Scandium.—P. T. Clève.—The source of this metal is gadolinite, in which it occurs in very small quantities. It probably does not belong to the yttrium group.

Russian Correspondence.—J. Lermontoff.

M. Gustavson reports on the compounds formed by cymol with aluminium chloride and bromide.

M. P. Goloubeff announces that the product of the reduction of nitrobenzol which he obtained in 1874 is identical with amido-desoxybenzoïn.

MM. A. Eltekoff and G. Lagermark have studied the action of sulphuric acid upon acetylen.

M. Potililtzine gave a preliminary communication on the action of oxygen upon haloid salts and the displacement of bromine by chlorine in anhydrous metallic bromides.

M. Louguinine described his results on the influence which the substitution of electro-positive and electro-negative elements, and also of the groups NO_2 and NH_2 for hydrogen, has upon the thermic phenomena accompanying the formation of salts. Aniline and para-toluydin possess very feeble basic powers compared with ammonia. Aniline, when combining with hydrochloric acid, evolves only 7.44 cal. The introduction of an atom of chlorine diminishes the liberation. The heat set free by nitro-para-toluydin in the same reaction is feebler again. In the substitution-products of the phenols the author finds that the introduction of the group NO_2 increases the liberation of heat on combination with Na_2O .

M. J. Goldstein has investigated the boiling-points of the saturated hydrocarbons of normal structure.

M. Andrianowsky publishes researches on the action of the acetic and sulphurous anhydrides upon aluminium chloride.

Notes.—M. S. M. Joergensen.—These relate to an anhydrous sodioferric pyrophosphate; a platinoso-platinic oxide, Pt_3O_4 ; and to the action of silver nitrate upon chloro-platinic acid.—*Journal für Praktische Chemie.*

Atti della R. Accademia dei Lincei.

Fasc. 5, April, 1879.

Influence of Boracic Acid upon Acetic Fermentation.—Prof. A. Herzin.—If an aqueous solution of boracic acid is added to pressed grapes in the proportion of one-twentieth by volume, fermentation is neither arrested nor retarded. The wine produced seems rather clearer than usual, but does not in any way betray the presence of boracic acid. But if the wine is placed in conditions most favourable for conversion into vinegar this change is absolutely frustrated. The author holds that the acetic micoderma lives at the expense of vinegar already generated and not of alcohol, and is a consequence rather than a cause of the chemical changes which are prevented by a trace of boracic acid.

Supposed Identity of Colombin and Limonin.—E. Paterno and A. Ogialoro.—Already noticed.

The Difficulty of obtaining Sulphuric Acid Perfectly Free from Arsenic and other Points Relative to Arsenic.—S. Sella.—For the detection of arsenic in sulphuric acid the author proposes to distill with lead chloride, testing the first portions which pass over with hydrogen sulphide. He suggests treatment with lead chloride for the final purification of sulphuric acid, partially freed from arsenic by sulphuretted hydrogen. Vitreous arsenious acid is not soluble in ponderable quantities in ether, methylic and amylic alcohols, and chloroform. Benzol, oil of turpentine, and the light petroleum oils do not take up a trace.

Fasc. 6, May, 1879.

New Method of Preparing Phenol-glycolic Acid and on Pyro-gallo-triglycolic Acid.—Dr. Piero Giacosa.

—The author heats together in the water-bath equivalent quantities of phenol and mono-chloroacetic acid, and adds little by little for each part of phenol 4 parts of a solution of caustic soda (sp. gr. 1.3), stirring the mixture well. The liquid boils, and while still hot forms a crystalline mass, which is the sodic salt of phenol-glycolic acid.

Researches on Boron and Vanadium.—Prof. Bechi.—The author has detected traces of boron in the marls and limestones of Montecatini, in Carrara marble, in basalts and lavas from Etna, and in trachytes from Tuscany. The same element has been discovered in the ashes of plants and in albumen extracted from the blood of oxen. He has found vanadium in argillaceous limestones, in shales, sands, and the ashes of plants.

Crystalline Form of certain Compounds of the Aromatic Series.—Dr. R. Panebianco.—Not capable of useful abstraction.

Chemiker Zeitung.

No. 23, June 5, 1879.

In this issue is a description of the Chemical Department of the Berlin Industrial Exhibition. Particular mention is made of the dyes displayed by the Aniline Company (Aktien Gesellschaft für Anilin-fabrikat), of Rummelsburg. Among their colours are eosin and methyl-eosin, coccin, mandarin, aurantia, silver-grey, malachite-green, Capri-blue, and phosphin.

J. K. Kothe exhibits an alcoholic distillate from pine-leaves, recommended as preferable to phenol for disinfecting sick-rooms.

Dr. Elsner, in replying to the recent strictures of Prof. W. Knop, complains that certain of the highest authorities in chemistry seem to discourage the war waged against sophistification.

The German import duty of 10s. per ton on crude iron took effect on June 30.

No. 24, June 12, 1879.

Complaints have arisen concerning the pollution of the Elbe by the waste waters of the Stassfurt and Aschersleben works.

Prof. C. T. Neubauer died at Wiesbaden on June 2.

The Grey Modification of Tin.—According to Schertehl, tin, under certain unknown circumstances, becomes so brittle as to be crushed between the finger-nails, and has the sp. gr. 5.8. If boiled in water it recovers its ordinary colour and texture, while the sp. gr. rises to 7.3.

According to the *Technologists* common rosin prevents the formation of acetic acid in fermented liquids without having any disturbing effect on the process of alcoholic fermentation. The peculiar effect of the hop may be due, it is suggested, to its resinous matter rather than to its oils. Resin is added to sweet wines in Greece.

Physiological Action of Atropin.—According to Petit atropin in doses of 0.030 grm. in the form of a subcutaneous injection produced no perceptible effect upon dogs.

No. 25, 1879.

The Berlin Exhibition.—Schmidt and Hänsch exhibit the extremely compendious and portable pocket spectroscope designed by Prof. Vogel. L. Reimann displays a balance which, when loaded with 500 grms., turns distinctly with 0.5 milligrm. P. Stuckrath has a small balance of aluminium carrying only 1 grm., but turning with 1.100 milligrm.

Source of Hippuric Acid in the Urine of the Herbivora.—The origin of this acid has long been ascribed to some substance present in hay, but which has not been identified. Loew, however, has found quinic acid to the extent of 6 grms. in 1 kilo. of air-dried hay.—*Journal Prakt. Chemis.*

Benzoic Acid in the Juice of Cranberries.—Loew has detected this acid in the juice of *Preisselbeeren* (the cranberries of northern and eastern Europe), and considers it the cause of their remarkable antizymotic and antiseptic power.

New Colouring Matter from Ortho-amido-phenol.—Aniline, according to Glaser and Schmitt, can be converted into azobenzol by potassium permanganate or chloride of lime. G. Fischer attempted in the same manner to convert ortho-amido-phenol into the corresponding azophenol. Instead of the latter compound he obtained a colouring matter which sublimed in garnet-red needles, but in very small quantity. On employing potassium ferricyanide as the oxidising agent he got a more plentiful yield. The new substance has slightly basic properties. It dissolves in benzol and in all acids, forming with the latter blue or deep violet solutions. Its formula is $C_{24}H_{10}N_3O_2$.—*Journal Prakt. Chemis.*

Nickelising without Electricity.—According to the *Moniteur Industriel* so much of a salt of nickel is introduced into a 5 or 10 per cent solution of zinc chloride till it assumes the ordinary colour of a nickel-bath. The articles to be coated, previously well cleaned, are laid in this solution and the process is complete in from half an hour to an hour. Cobalt can be deposited in the same manner.

Diamond Hunting in China.—Chinese diamonds are chiefly brought from the province of Shantung. Men put on thick shoes of straw and simply roam about the valleys and the rivers. The rough and pointed diamonds penetrate into the straw and stick fast. The shoes are finally collected together in heaps and burnt, when the diamonds remain in the ashes.—*Technologiste.*

Universal Stomach Powder.—Prince F. W. Barella, of Berlin, appears to be the proprietor of this medicine, which consists of 92.70 per cent bicarbonate of soda, 4.00 per cent common salt, and 2.30 per cent carbonate of lime. A box containing 100 grms. costs eighteen pence.

Colouring Matters from Coal.—Dr. Meusel, of Breslau, prepares colouring matters by treating coal-dust with nitric acid, or with a mixture of alkaline nitrates

and sulphuric acid, or with other oxidising agents. A part of the coal is thus rendered soluble with a deep brown colour in alkaline hydrates or carbonates. The black residue serves as a body colour. The brown alkaline solution can either be used at once, or metallic salts of a black or dark brown colour may be precipitated from it, or the precipitate produced by the addition of acids may be used as a pigment.—*Die Allg. Polyt. Zeitung.*

Bleaching Ostrich Feathers.—According to the *Moniteur des Produits Chimiques* feathers are bleached in a bath of 10 grms. barium peroxide to 1 litre water, heated to 30°. In this they remain for forty-eight hours, and are then washed, treated with weak hydrochloric acid, and dried.

No. 26, 1879.

Anthocerin.—F. v. Müller.—This compound is a volatile alkaloid obtained from the aqueous extract of an Australian Solanaceous plant, *Anthocoris viscosa*. The author will shortly report on its composition and on its physiological action.—*Zeitschr. Oest. Apoth. Ver.*

A Mordant from Lees of Wine.—Fresh green lees, with the addition of two-fifths sodium tartrate, are evaporated down to one-sixth the original volume. 15 grms. of Cologne glue and 10 grms. tannic acid are added. The mass is pressed, rubbed over with alcohol and tannic acid, dried in the air, and powdered. For use it is to be further mixed with one-fifth per cent sodium tartrate. It is recommended for dyeing full shades in wool and silk, and is said to render the aniline colours permanent. In dyeing woollen cloth a decoction of Saponaria root is added both to the mordant and the dye-bath.

Action of Vinegar upon Alloys of Lead and Tin.—R. Weber.—Vinegar attacks pure tin as well as alloys with lead, the quantity of metal dissolved increasing with the proportion of lead present. Alloys of tin and lead, to which 4 per cent antimony had been added, were also attacked, and lead entered into solution.

Mineral Wealth of Turkey.—The following metals are entirely absent:—Tin, cobalt, nickel, bismuth, and uranium. Chrome iron ore, emery, and copper are plentiful. Coal is found only in the basin of Ereğli and Amastra on the Black Sea. There are numerous petroleum wells on the Persian frontier.

No. 27, 1879.

Action of Chloride of Lime upon Ethylic Alcohol.—The formation of chloroform by this reaction and the transition thus effected from the ethyl to the methyl series have not yet been explained. Schmitt and Goldberg find that if chloride of lime is added to absolute alcohol heat is generated in from seven to ten minutes, and distillation sets in. Along with much alcohol a yellowish green oil passes over, which on exposure to light or heat is decomposed with explosion, giving off hydrochloric and hypochlorous acids. In the distillate there are found after the explosion mono-chlor-acetal, dichlor-acetal, chlorinised methyl-ethyl-ether, alcohol, aldehyd, and chloroform.—*Journal Prakt. Chemis.*, xix., 393.

Tests for Oils.—Maumené considers that the only trustworthy characteristic of oils is the heat liberated on mixing 50 grms. of the oil with 10 c.c. concentrated sulphuric acid, which is added by means of a pipette, and the whole then well stirred with a thermometer for some minutes, the highest temperature being read off. With pure olive oil the rise is 42°, whilst with linseed it amounts to 103°.

Reimann's Färber Zeitung,
No. 25, 1879.

This issue contains nothing of general interest.

No. 26, 1879.

Benzyl Blue.—This new dye is probably a product of the substitution of rosanilin by three atoms of benzyl. It is readily soluble in boiling water, and is applicable both to animal and vegetable fibres.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 10, July 3, 1879.

This issue contains no original chemical matter.

*Verhandlungen des Vereins zur Beforderung des
Gewerbfleisses.* May, 1879.

Aniline-black.—Dr. R. Kayser.—The author has examined the aniline-blacks produced by means of ammonium vanadate, copper sulphate, and free hydrochloric acid. He recommends that such aniline oils as are free from toluoydin, *i.e.*, the lightest, should be exclusively used for aniline-blacks. He rejects Nietzki's view of two isomeric bases, the one of a more blue or violet, and the other of a brownish cast.

Aniline-black.—Dr. R. Nietzki.—The author considers that aniline-black is probably closely connected with the indulins, its colour in a basic state being violet, but when acid, of a greenish cast.

La Correspondance Scientifique.
No. 45, June 10, 1879.

This issue contains no original chemical or physical matter.

No. 46, June 17, 1879.

The Mine-Explosion at Frameries.—It appears from the report of Prof. Burat that more than 100,000 cubic metres of fire-damp must have been evolved in a very short time. All the 200 safety lamps (Mueseler's) were extinguished without causing ignition. The gas issuing from the shaft caught fire *outside* the mine (possibly from the engine furnace), and burnt with an enormous flame. When the supply was becoming exhausted the burning gas ran back into the interior of the mine, followed in its retreat by atmospheric air, and occasioned nine successive explosions. The whole occurrence is considered unexampled in the annals of coal-mining.

Priority of the Discovery of Artificial Ultramarine.—E. Guimet produces documentary evidence that his father, J. B. Guimet, had manufactured artificial ultramarine as early as 1826, and had sent samples to different artists, who pronounced it of excellent quality. In 1827, Ingres used some of it in the decoration of the museum. In 1828, at the time of Gmelin's visit to Paris, and of his conversation with Gay-Lussac, it was already an article of commerce, and had earned the approval of eminent artists.

Die Chemische Industrie.
No. 6, June, 1879.

The Tar Colours and the Electric Light.—Dr. Greiff.—The author contends that even supposing the gas manufacture should ultimately be abandoned the tar colours could be prepared from the residues left on rectifying the petroleum of the regions on the Caspian. These are estimated at 120 million kilos. yearly, and are ten times richer in benzol and five times richer in anthracene than is coal-tar. The American petroleum has not yet been examined from this point of view, but it will probably also prove to be a rich source of aromatic compounds.

MISCELLANEOUS.

A Chemical Reminiscence (Reply to "S.")—

Sankey brook cannot hold its wealth
Which it sadly bears to the sea far out.
If you knew the bait to catch its fish
You would never lament its want of trout.

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THE CHEMICAL NEWS.

VOL. XL. No. 1027.

BEHAVIOUR OF CHLORINE AT A HIGH TEMPERATURE, OR RESULTS OF VICTOR MEYER'S RECENT RESEARCHES.*

By WATSON SMITH, F.C.S., F.I.C.

IN the course of their determinations of the vapour-densities of inorganic substances at high temperatures in the now well-known apparatus devised and perfected by them, my friends Professor Victor Meyer and Carl Meyer have recently turned their attention to the estimation of the vapour-density of chlorine at various high temperatures, viz., the following:—620°, 808°, 1028°, 1242°, 1392°, 1567°. The higher of these temperatures, determined by calorimetric observations, are approximate merely. The chlorine was prepared absolutely pure by heating PtCl_2 (or Pt_2Cl_4), and was thoroughly well dried. The resulting vapour-density observations at the named temperatures were as follows:—

	Found.		Theory.
	I.	II.	
At 620° ..	2.42	2.46	For $\text{Cl}_2 = 2.45$ (vapour density.)
808 ..	2.21	2.19	
1028 ..	1.85	1.89	
1242 ..	1.65	1.66	
1392 ..	1.66	1.67	
1567 ..	1.60	1.62	
			1.63 is exactly two-thirds of 2.45

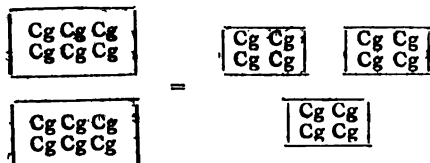
Thus, at 808° an evident "splitting-up," or decomposition, or dissociation commences, which at 1227°, 1392°, and 1567° becomes and remains constant. The molecular weight, then, of chlorine at from 1242° to 1567° would appear to be—

$$\frac{2}{3} \times 71 = 47.3.$$

With iodine quite analogous results were obtained, but the experiments are not as yet concluded in this direction. Bringing these observations to bear upon chlorine still regarded as an element, and regarding the amount of permanent expansion resulting in the contraction of vapour-density from 2.45 to 1.63 (viz., two-thirds), also bearing in mind the fact that chlorine in combining with hydrogen to form hydrochloric acid gas does so volume for volume (in equal volumes), uniting without condensation—then it becomes evident that (1) the chlorine has suffered a molecular "splitting-up" or dissociation; and (2) the chlorine molecule, $\text{Cl} \dots \text{Cl}$ cannot even be regarded as consisting of 3 atoms, but must at least contain 6 atoms. Hence the weight of an atom of chlorine would be derived as follows from the molecular weight 70.8—

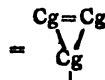
$$\frac{70.8}{6} = 11.8, = \frac{35.4}{3}.$$

The decomposition may then be represented as taking place in the following manner:—

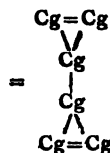


* Report from notes taken at the meeting of the Chemical Society of Zürich, held Monday, July 21, 1879, during V. Meyer's lecture.

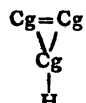
Therefore $\text{Cl} = \text{Cg}_3$, and $\text{Cl}_2 = \text{Cg}_6$, where Cg stands for the hypothetic atom, *Chlorogene*, having the weight 11.8. Further, the atom of chlorogene, Cg, is probably trivalent, and Cl then =



Cl_2 (the molecule) =



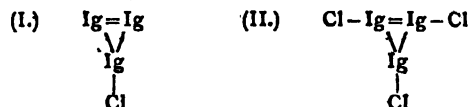
And hydrochloric acid—



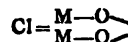
Also iodine, I =



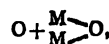
where Ig represents the hypothetic atom Iogene. Thus, the formation and constitution of the iodine chlorides would be thus explained:—



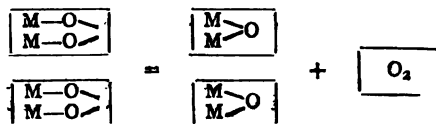
Prof. Meyer next proceeded to a practical examination of the old—once celebrated, but now nearly forgotten—"Murium Theory" of Berzelius, according to which it was supposed that chlorine was, in fact, an oxygen-containing substance—an oxygen compound. This research he has carried out in conjunction with Carl Meyer, and with the object of discovering if any basis existed for this old theory. The experiments have been conducted with the utmost exactitude and care, and every possible error kept away, as far as can be seen—in fact Victor Meyer has invited with gratitude any criticism, and still continues to do so, bearing upon the methods adopted and mode of conducting the experiments. He shows, for example, that the porcelain containing vessel (placed in a Perrot's furnace, and through which the chlorine gas was passed) could not have contributed any oxygen. Further, the chlorine was dried with every precaution, by passing it over a layer of phosphoric pentoxide 1 metre in length, &c. Experimental evidence appears to indicate beyond a doubt that to all appearance ("Möglicherweise" is the word Victor Meyer wishes at present to be used) by heating chlorine oxygen can be obtained. This being proved, then the splitting up of chlorine at a high temperature will give rise to the following probable theory:—If—



where M = Murium, then this atom or Cl, at the high temperature mentioned, yields—



and the molecule $\text{Cl} \dots \text{Cl}$ is decomposed as follows:—



Thus ends the matter so far, but I need scarcely add that Victor Meyer's researches are continuing, and he is prosecuting them with unabated vigour, both in the directions of most stringent confirmation and further advance. For myself, since hearing of the discovery, I have thought much of and pondered over the wonderful relationship as to melting- and boiling-points calculated from the absolute zero, for the first time observed by Dr. T. Carnelly and W. C. Williams, of the Owens College, between chlorine, bromine, iodine, and sulphur, selenium, and tellurium (see CHEMICAL NEWS, vol. xxxix., p. 286). On consideration of this relationship, taken now in conjunction with Victor Meyer's great discovery, it seems to me very probable that all the six named, so-called, elements contain one and the same essence ("Gründstoff" is the more exact term in the German), and will be ultimately decomposed so as to yield this essence common to all six (Cl, Br, I, S, Se, and Te). This is not a hap-hazard "phantasie," but still further founded on, and suggested by, a consideration (combined with the foregoing one) of the analogy afforded by the melting- and boiling-points of the organic hydrocarbons and other carbon compounds arranged in certain series, viz., homologous, polymeric, and isomeric series.

These compound bodies contain a common so-called elementary body, carbon, combined with hydrogen in certain definite proportions, so that definitely constructed or ascending series may be arranged, in which melting-points and boiling-points are found also of definite arrangement, ascent or descent. In such compounds, too, the hydrogen may be replaced by the so-called elements, chlorine or bromine, and still furnish respective series of the kind named. The hydrogen in CH_4 , too, may be considered replaced by sulphur, giving the compounds CH_3S and CS_2 . Again, a mixed compound of S and Cl with C in the same molecule may be formed; e.g., by the action of dry chlorine on carbon disulphide, $\text{CS}_2 + \text{Cl}_2 = \text{SCl}_2 + \text{CSCl}_2$. Again, besides the combination of iodine with chlorine, a compound of iodine with sulphur is known, I_2S_2 , also of bromine and sulphur. Of chlorine with sulphur two compounds occur, S_2Cl_2 , boiling at 136° nearly, and SCl_2 boiling at 164° .

It does not seem, then, at all impossible that the chlorine group may furnish a kind of homologous, perhaps polymeric, series, and the sulphur group another homologous series, derivable from the first by some simple means, similar to those applicable in the hydrocarbon series. In other words, that a species of polymeric relationship possibly exists between the chlorine group members, and another and higher, more complicated, kind of polymerism connects this group with the sulphur group, and connects it intimately. To take a case of polymerism in a hydrocarbon group observing boiling-point relations, the group given by Schorlemmer in his "Chemistry of the Carbon Compounds" (p. 35) deserves notice. This group comprises the following:—

Acetylene,	C_2H_2	
Benzene	C_6H_6 , and adding boiling-pnts. = 353° .	
Styrolene,	C_8H_8 , reckoned above -273° as zero	$= 419^\circ$
		$= 487^\circ$.
Hydonaphthalene, $\text{C}_{10}\text{H}_{10}$		

The ordinary homologous series with their gradually rising boiling-points are too well known to need further than mentioning.

Here it may be noticed that the boiling-point of styrolene is nearly the mean of those of the other two members, benzene and hydonaphthalene, standing next above and below it; just as in the case of chlorine, bromine, and iodine, whose boiling-points are respectively 240° , 318° , and 473° (reckoned above absolute zero by Carnelly and Williams). 318° for bromine is not very far from the mean between (Cl) 240° and (I) 473° , which is 356° . All this, of course, is a speculation intended to illustrate the line of analogy suggested by the comparisons mentioned; still future experimental research may show that there is truth in it, but the patient experimental work alone can

give the means of clearing up the mist which at present seems to overhang the whole question. It has often required the exertions of a master-mind and a master-hand to convince men that *they really are in a mist*, and the researches of Victor Meyer have recently done much, one may indeed say *most*, to convince the world of chemists that their elemental atmosphere is by no means so clear or cloudless as they had once imagined; also that much requires yet to be done ere the clear truth-light shines down upon them with unsullied brightness and splendour!

ON MANURE PHOSPHATES.

By K. WALTER,
Chemical Engineer, Aurelais, Belgium.

(Concluded from page 38.)

To the third group, the phosphates which are quite or for the most part soluble in citrate of ammonia belong some few sorts of guano; those, for instance, which contain up to 30 per cent of their phosphoric acid soluble in citrate. There are also some few natural phosphates containing a few per cents of their phosphoric acid soluble in the citrate. Then the so-called retrograded superphosphates. I have had myself superphosphates in work which held in the beginning 11.5 per cent of phosphoric acid soluble in water; after three years they only contained 1.5 per cent of it. It is a great injustice against superphosphate manufacturers that in England as well as in Germany this retrograded phosphoric acid is counted worthless in the market; the more so since for more than ten years an exact process of analysis has been known which determines the difference between natural and retrograded phosphoric acid. There exist lots of superphosphates which contain, after four to five months' storing, one-tenth to one-sixth of their phosphoric acid formerly soluble in water in a retrograded state, which latter is soluble in citrate of ammonia. Different raw phosphates have not been employed up to the present time because of their property of forming superphosphates which retrograde. The precipitated phosphates originating from a well conducted manufactory chiefly belong to this group. Such a one has 85 to 95 per cent of its phosphoric acid soluble in the citrate; it has not, however, up to the present time been sold either in Germany or in England, at all events not at the price it is worth.

Toulie years ago proved, by very extensive experiments (see his work mentioned on page 37), that bibasic phosphate of lime (which is soluble in the citrate) has the same effect as superphosphate, even in poor sandy soil. Alas! neither the English or German agricultural chemists would credit this experience—and yet they were unable to give the slightest proof to the contrary. Even when Toulie's experiments were shown to be quite correct by their being repeated by Dr. Petermann, and after the latter officially announced (in the year 1877) that from January 1, 1878, in all phosphates coming under the control of the Belgian Agricultural Stations all phosphoric acid soluble in citrate of ammonia must be counted as assimilable and of the same value as soluble in water, not a hand in England or Germany was lifted to follow the praiseworthy example.

Likewise the trials of Grandeau* (manager of the Chemical Agricultural Station of Nancy, France), on a very extensive scale and repeated for several years, have proved that precipitated acts not only with the same speedy efficacy as superphosphates, but even with greater.

It is further shown, by the experiments of Petermann, that phosphate of iron and of alumina (soluble in citrate) are even far more efficient when employed as manure phosphates than even superphosphate and precipitated

* "Annales de la Station Agronomique de l'Est, France," vol. ii. (See also Biedermann's *Centralblatt für Agricultur Chemie*, 1876, Band 2, Seite 650.)

phosphate of lime. Up to the present time it has been taken for granted by nearly all chemists that the process which takes place in some sorts of superphosphates, commonly called retrograding, is caused by oxides of iron and alumina, which form, with a part of the mono-basic phosphate of lime, bibasic phosphates of iron and alumina insoluble in water. Be it so, there is then an additional reason to value the retrograded phosphoric acid as not only equal but even higher than the superphosphate itself. If *superphosphate* comes into contact with the *soil* this process does not seem to take place, but rather that *tribasic* phosphate of lime, and perhaps of iron also, is formed. For if bibasic phosphate of iron and alumina is employed directly for manuring, the result is far more favourable than with precipitated phosphate (soluble in citrate), the result of this latter being already superior to that of superphosphate.

These are facts stated by different eminent agricultural chemists, and put to trial by the experience of years and years. It is not the case, however, that only bibasic phosphate of lime, iron, and alumina are soluble in citrate of ammonia, the tribasic are also soluble under certain circumstances. Nobody, I think, would be able to make a precipitate (dried at not over 100° Celsius), of which not always a certain portion is soluble in the citrate, at all events I myself never succeeded. On the contrary, it is very difficult to make a product which is thoroughly soluble in the citrate.

I have worked for three years at this subject, but I must confess the results obtained in the laboratory differ from those in the manufactory, a result which I cannot as yet explain. It would occupy too much space to enter into all the reactions which can occur in phosphate making. True it is that for the chemists the analysis after the citrate method is very tedious, but this is not a sufficient reason to suppress a progress made by science and proved by practice; and if the French and Belgian chemists were obliged to accommodate themselves to it, the English and German chemists will, sooner or later, be obliged to do the same.

The manufacture of precipitate is only carried on with advantage in places where muriatic acid, and at the same time mineral phosphate, are to be had at a very cheap rate. There are at present only two works existing in France, and more will hardly be erected, the muriatic acid having too great a value in France. One of these works is carried on with great success, being able to reckon (an exceptional case) the muriatic acid at an extremely cheap rate, and being in the neighbourhood of rich layers of phosphate containing very little iron, alumina, and carbonate. Besides this, the work is managed extremely well, having the advantage of the many years experience. The other work mentioned has not yet made a product sufficiently rich in phosphate soluble in citrate. This branch of manufacture is only remunerating when the product has at least nine-tenths of its phosphoric acid soluble in the citrate.

In Belgium there exists one work for manufacture of precipitate, erected by the writer. The delivery of muriatic acid at a cheap rate being assured for several years, and the raw phosphates being cheap enough, the work can be carried on with fair success. The raw phosphate is coprolites from the neighbouring French Ardennes, containing 35 to 40 per cent of phosphate of lime, and only 7 to 8 per cent of carbonate. The phosphate of Mons, which would be for here so very convenient, is, as mentioned already, not fit to be used for this manufacture; it wants more than double the quantity of muriatic acid for its decomposition than it ought to do for the manufacture to be profitable.

In most cases, phosphates which contain much phosphate of alumina and iron are also unfit for the manufacture. This is the reason why the work erected at Mühlheim on the Rhine, Germany, by an English company did not succeed. They intended to work Nassau phosphate, which is not fit for the manufacture—containing too much

iron and leaving too much phosphoric acid in the residues; besides this, muriatic acid can be sold in Germany at a much higher rate than it could be employed with profit for the manufacture in question. Then, also, the management was not at all up to the task of making soluble phosphate. It is the problem of the precipitate manufacture to employ phosphates which are too poor for the superphosphate manufacture.

In Belgium and France the same rate is realised for the precipitate as for the superphosphate—0.80 fr. per kilogram. of phosphoric acid, taken in large quantities; the former of course pays more. There is no doubt the time will come that good precipitate will be paid for at even a higher rate than superphosphate, if only on account of its richness, which renders it more fit for transportation to a distance. Already more than half the quantity produced is sent to the French and Dutch Colonies.

Superphosphate with 20 per cent of phosphoric acid is not easily to be found in the market, while precipitate can easily be made with 35 to 38 per cent of phosphoric acid. Because of its finer division it is also much better adapted for top-dressing (a method of manuring which in later years has been more and more employed), not to speak of its harmlessness: while superphosphate before decomposed by the soil is a poison to the young plants.

To work in the direction which Petermann and other agricultural chemists' results of experiments indicate, it would be very desirable to bring into the precipitate as much phosphate of iron and alumina as possible; but the technical difficulties have been up to the present too great. Not only are large quantities of phosphate of iron and alumina, insoluble in the citrate, formed in the process of precipitation, but precipitate with too much of the latter phosphate is, on account of its fine division, too difficult to work in the filters and presses. If a precipitate contains one-fourth of its phosphoric acid united to iron and alumina, this is the utmost obtainable until other means are found.

To the last of the four groups belong the superphosphates. They are manufactured of from 11 to 12 per cent of phosphoric acid soluble in water. One of less than 11 per cent is hardly to be found now in the market; and to make them richer than 22 per cent, and at the same time dry, will be also too costly. As mentioned already, in Belgium and in France the phosphoric acid soluble in water is no longer considered in the analysis; all that is soluble in citrate of ammonia is taken for soluble and immediately assimilable. It is my opinion the experiments of Dr. Petermann and others, at present going on, will prove evidence—as already, however, proved practically—that even certain phosphates, *not soluble in citrate of ammonia* are just as speedily assimilable as the before-mentioned.

Superphosphate is, strictly taken, a necessary evil. No one will be hardy enough to maintain that monobasic phosphate of lime (superphosphate) is assimilated by the plant as such; and every one who is experienced in the subject will agree that the superphosphate finds in the soil more carbonate of lime, oxides of iron and alumina, than necessary to transform it speedily into bibasic and tribasic phosphates. The two latter are, then, taken up and put into an assimilable state by the salts of ammonia, soda, potash, and the carbonic acid to be in such solution assimilated by the plants.

It is equally a great question if the arsenic brought into the soil by means of the superphosphate in the long run does not create an injurious action. Sulphuric acid employed for superphosphate making contains sometimes quite formidable quantities of arsenic, which in this manner is incorporated with the soil in a soluble state. However, natural phosphates acting mostly too slow, bones and guano not being available in the market in sufficient quantity or sufficiently cheap, and as advantageous means of producing sufficient quantities of lime, iron, and alumina phosphates are not yet known, the superphosphate will still, therefore command the market. The manufacture of pre-

cipitate will go on beside the superphosphate manufacture, like the soda-ammoniac process beside the old process of making soda-ash.

There is still a product to mention which is made in Germany, liquid phosphoric acid, sold in barrels, of course impure, and not free from monobasic lime and iron phosphate. It is used for certain mixed manures, which have to be comparatively rich in phosphoric acid.

GENERAL OUTLINE OF A NEW THEORY OF MOLECULAR SUBSTITUTION, AS CONCEIVED AND ELABORATED FROM THE STANDPOINT OF THE "TYPO-NUCLEUS" THEORY.

By OTTO RICHTER, Ph.D.

IN pondering the precise nature of the various molecular changes which accompany the process of substitution proper I was not long in satisfying myself that, over and above the method of reciprocal or two-sided transfer which characterises the ordinary process of double decomposition (CHEMICAL NEWS, vol. xxxix., p. 59), other modes of transmission were yet resorted to in nature's laboratory which had hitherto eluded the attention of chemists, either because the accompanying changes were strictly intramolecular, or because they consisted in the one-sided transfer of one single subjunct only. It is with the aid of these broader views and conceptions that I hope to succeed in expounding the true *modus operandi* of that wonderful species of mechanism which presides over this particular order of chemical phenomena. From an extensive series of researches on this subject I have, further, been led to conclude, that the said mechanism is indebted for its main source of motive power to the fundamental law of quantivalence or thermal bond-relationship, which again, for its proper manifestation, depends entirely on the support and co-operation of the fundamental law of multivalence or physical impact-relationship. The *nexus causalis* thus affirmed to exist between these three kinds of agencies naturally suggested to me the exact logical order in which the two laws just spoken of ought to be studied and discussed. The present communication bears the title:—

On the Fundamental Law of Multivalence or Physical Impact-Relationship.

In reflecting on the primary cause of those simple numerical relations which are found to obtain between the equivalent volumes and vapour-densities of æriform bodies, I have at length been brought to recognise the general law which dominates this section of natural philosophy. The law in question, is regarded by me as the outcome of certain dynamic relations established between the elasticity, vapour-density, and heat-emitting energy of æriform substances, and may be briefly enunciated as follows:—*Under the same conditions of temperature and pressure equal volumes of æriform bodies, whatever their atomic weight and chemical composition may be, always contain an equal number of molecules.* Taking the reality of this law for granted my next duty will be to expound the fundamental law of multivalence, and to point out its intimate connection with the law of æriform bodies just enunciated. In my opinion the law of multivalence is the outward manifestation of a powerful tendency inherent in all chemically simple bodies to undergo the process of self-condensation, in virtue of which two or more univalent molecules, $2E_1$, may by a species of physical impact become cemented together in such a way as to give rise to a series of progressively, more complex, and specifically heavier molecules, for which I shall employ the terms—bivalent, trivalent, quadrivalent, &c. The class of molecules under consideration have, moreover, this characteristic feature in common, that they are

capable of preserving the fundamental type originally impressed upon their parent molecules. In further illustration of this subject I shall now present the reader with a brief summary of such valuable experimental evidence as may be adduced in proof of the general soundness of this new doctrine of molecular self-condensation.

Commencing with the class of non-metallic elements we encounter in the highest member, $2S_{12}$ of the sulphur series, $2S_2$; $2S_4$; $2S_6$... $2S_{12}$, a notable instance of self-condensing energy. To this molecule with a vapour-density six times greater than what by theory pertains to the first or parent molecule of the series, there became added afterwards a second modification, $2S_8$, which, by its vapour-density, is evidently pointing to the second member of the series. A similar series, $2O_2$; $2O_4$; $2O_6$, but of half the length only, is met with in the case of oxygen, the second member of which corresponds no doubt to the ordinary atmospheric variety, while the third member, so-called ozone, is a more recent discovery. Turning to the element nitrogen only one modification is familiarly known to us, but in striking contrast with the common practice I have good reason for identifying it with the second member, $2N_4$ ($N_2=14$), of the series. It is worthy of mention that, although the first or generating members of the three series before us have not yet been obtained in a state of isolation, they are, nevertheless, by common consent and from purely theoretical considerations only, believed to exist as such in a great variety of chemical combinations. Glancing, in the next place, at the class of metallic elements I shall select from among their more volatilisable members, first, the quasi-metallic element, phosphorus, the only known modification of which is by its vapour-density held to correspond to the fourth member, $2P_8$, of the series, while the lower members can be shown to exist as such in combination with sulphur, oxygen, and other non-metallic elements; secondly, the truly metallic element, mercury, which belongs to the rare class of metals, whose generating molecules in passing from the liquid into the æriform state, so far from undergoing a process of self-condensation, are actually, to judge from their vapour-density, made to split up into two equal halves, so that in the case of mercury the resulting semi-valent molecule will claim to be expressed by the symbol, $2Hg$ ($Hg_2=400$). We are now in our general survey brought face to face with hydrogen, the specifically highest and certainly the most remarkable of all the metallic elements. I will only observe here that in the free state hydrogen seems to be entirely destitute of the power of self-condensation, whereas after being combined with carbon, whereby it gives rise to the prolific family of the hydrocarbons, that same element appears to have acquired a degree of self-condensing energy which places it on a par with sulphur, carbon, and other elements.

Having now briefly explained to the reader the general import and significance of the law of multivalence as applied to chemically simple substances, whereby it gives rise to a number of variously extended series of derivatives, which have all of them this characteristic feature in common, that they are constructed on the fundamental type of principal nuclei, I shall in the next place consider the law of multivalence in connection with that class of chemical combinations which are formed by the direct union of the aforesaid principal nuclei with outer conjunct molecules (CHEMICAL NEWS, vol. xxxix., p. 48).

Confining my remarks to the four elements, carbon, sulphur, nitrogen, and oxygen, and to the first two series of derivatives, the following combinations have been selected and embodied in the annexed table of chemical formulæ.

In commenting on this scheme (next page) I may observe that the compounds before us belong each and all to the class of so called anhydrides, but although bearing a strong resemblance to each other in their general chemical character and deportment, I have satisfied myself that there exist certain important constitutional differences between them which it becomes incumbent upon me to

Table of Chemical Formulae Representing Combinations of Principal Uni-orbivalent Nuclei with one Species of Outer Conjunctions.

Univalent Nuclei.	Outer Conjunctions.	Bivalent Nuclei.	Outer Conjunctions.
1st Series $2C_2$ $2S_2$ $2N_2$	$\left. \begin{array}{l} O_2; O_4 \\ O_2; O_4; O_6 \\ O_2; O_4 \end{array} \right\} \left. \begin{array}{l} N_2 \\ N_2; N_4 \\ \text{vacat} \end{array} \right\} \left. \begin{array}{l} S_2; S_4 \\ \text{vacat} \\ \text{vacat} \end{array} \right\}$	2nd Series $2C_4$ $2S_4$ $2N_4$	$\left. \begin{array}{l} O_2; O_4; O_6; O_8 \\ O_2; O_4; O_6 \dots O_{12} \\ O_2; O_4; O_6 \dots O_{10} \end{array} \right\} \left. \begin{array}{l} N_2; N_4 \\ N_2; N_4 \dots N_8 \\ \text{vacat} \end{array} \right\} \left. \begin{array}{l} S_2; S_4; S_6; S_8 \\ \text{vacat} \\ \text{vacat} \end{array} \right\}$

point out. For this purpose I have bethought myself of dividing these anhydrides into two separate classes. Among the first class I shall include all those compounds whose outer conjunctions are composed of an even number of univalent sulphur, nitrogen, or oxygen molecules, and among the second class I shall include all those compounds whose outer conjunctions are composed of an uneven number of the aforesaid molecules. By this proceeding we are enabled to perceive at a glance the true reason why, as a general rule, the vapour-densities of the first class of molecules correspond to one-half only of their calculated value, while the vapour-densities of the second class correspond to the full amount of that value. Now, in my mode of viewing, the cause of this singular discrepancy must be sought in the circumstance that the univalent molecule of sulphur, nitrogen, and oxygen, and consequently also their multiples by an uneven number, stoutly refuse being split up into two equal halves, while their bivalent modifications seem to offer no formidable obstacle to that operation. On these premises we may readily trace the molecular changes which accompany, for instance, the splitting up of a molecule of anhydrous oxalic acid (dicarbo-trioxide), $2C_4O_6$, into a molecule of carboxide and a molecule of carbo-dioxide. Accordingly the first stage of the process, which requires the co-operation of two molecules of dicarbo-trioxide, will be marked by the transfer of a univalent molecule of oxygen from one molecule of the trioxide to the other, with formation of a molecule of dicarbo-dioxide, $2C_4O_4$, and a molecule of dicarbo-tetroxide, $2C_4O_8$, both of which, according to my theory, ought to be, as they are in fact, very prone to split up into two equal halves, namely, the former into two molecules of carboxide, $2C_2O_2$, and the latter into two molecules of carbo-dioxide, $2C_2O_4$. Applying our hypothesis to other combinations we can, further, understand why the vapour-density of hydroxide corresponds to the formula $2H_2O_2$, while the vapour-density of hydrochloride corresponds to the formula $2HCl$; for in either case the direct union of a univalent molecule of oxygen, on the one hand, and a univalent molecule of chlorine on the other hand, with a hydrogen nucleus gives rise first of all to the molecules $2H_2O_2$ and $2H_2Cl_2$, but owing to the circumstance that oxygen belongs to the class of indivisible elements, while both hydrogen and chlorine belong to the class of readily divisible elements, the final result will be that the molecule of hydroxide remains intact, while the molecule of hydrochloride splits up into two equal halves. In connection with this subject I may yet be permitted to dwell on the seemingly anomalous behaviour of certain compounds, which, from the bivalent character of their principal nuclei, as well as their outer conjunctions, ought to prove by their vapour-densities that in obedience to the general rule they have undergone the operation of splitting, whereas in reality no such change has taken place, or at all events not within the limits of a certain low temperature. The first of these exceptions is cyanogen, which, by its vapour-density, claims to be represented by the formula $2C_4N_4$; nevertheless, and although it is composed of a bivalent molecule of carbon and a bivalent molecule of nitrogen, our substance refuses to split up into two equal halves, even at a high temperature. But apart from this there can be no doubt, on the other hand, that the hypothetical semi-cyanogen with the formula $2C_2N_2$, does really exist as such in a vast number and diversity of organic compounds, as, for instance, in aceto-nitrile, $2H_3C_2$; $2C_2N_2$, where it is formed by the action of ammonia on hydric acetate. The second exception is nitrous acid, which exhibits a curious anomaly

in this respect, that, below $82^\circ F.$, it possesses a vapour-density corresponding to the formula $2N_4O_8$, while at a higher temperature it shows a vapour-density corresponding to the formula $2N_2O_4$, clearly proving thereby that the nitrous acid molecule has under the influence of heat been made to split up into two equal halves.

In the preceding pages it has been my endeavour to familiarise the reader with the most striking and characteristic features of the law of multivalence, and to demonstrate by a great variety of strong experimental evidence its intimate connection with the law of vapour-densities. I will only add in conclusion that in my treatment of the subject before us I have as much as possible confined myself to what I considered absolutely necessary for the proper comprehension of the fundamental law of quantivalence.

COAL-TAR COLOURS AND THE ELECTRIC LIGHT.

By Dr. GREIFF.

THE author seeks to show that even in case the manufacture of gas were to cease, the supply of raw material for the production of artificial colours would not be imperilled.

In order to find a standard for the quantities of anthracen and benzol now required, it may be assumed that 30,000 kilos. of alizarin paste at 10 per cent are produced daily. The minimum quantity of pure anthracen needful would therefore be 3000 kilos. daily, or 900,000 kilos. per year. This necessitates the annual distillation of 360 million kilos. of tar, containing on an average $\frac{1}{2}$ per cent. of anthracen. The production of aniline amounts to 12,000 kilos. daily, requiring 13,300 kilos. benzol, and also obtained from the same 360 million kilos. of tar.

Future inventions will further heighten the importance of coal-tar so long as it is regarded as the sole source of the aromatic hydrocarbons.

But in case a partial scarcity of this substance should arise, we have firstly the proposal to obtain tar, not as a residue, but as a primary product by the dry distillation of coal, and to attempt the proportionate increase of its more valuable ingredients. A quantity of tar might be obtained by an improved construction of certain furnaces for metallurgical purposes, and of coke-ovens—for which, while the present ample supply is obtained from the gas-works, there is little inducement.

It is also possible that the worthless hydrocarbons in tar may be transformed into benzol and anthracen. The valuable products of coal-tar do not exceed 3 to 4 per cent, along with 30 to 40 per cent of oils, boiling at high temperatures, and of solids, a minority of which only are recognised as definite chemical individuals.

But there is a still simpler and easier method of obtaining aromatic hydrocarbons in more than sufficient quantity. Liebermann and Burg, Wichelhaus and Salzmänn, and other chemists, have passed wood-tar and the tar of lignite through ignited tubes, and have invariably recognised the formation of aromatic hydrocarbons. These observations would have a mere scientific value did they not extend equally to the mineral oils of the Caspian and the tar—at present useless—which is obtained as by-product from their rectification. In order to decide

this question we must consider whether the raw material is present in sufficient quantity; what is the yield of benzol and anthracen; and finally, what is the cost of the process? All these points receive a favourable reply. Letny (*Dingler's Polytech. Journal*, 229, p. 353), shows that the product of crude mineral oil for the year 1875 was 240 million kilos.—a quantity easily susceptible of being greatly multiplied. After obtaining the lumirous and the lubricating oils, more than one-half remains behind, and is partly used as fuel on board the steamers on the Wolga. Letny has decomposed this residue at a red heat and obtained from it a product, about one-third of the original weight of the tar, which yields 4.6 per cent benzol, 5.2 toluol, and 3.1 per cent of anthracen, and which can be worked far more easily than coal-tar, though yielding the same hydrocarbons in the same succession. The 120 million kilos. of tar represent, therefore, as far as the yield of valuable aromatic hydrocarbons is concerned, from 200 to 400 million kilos. of coal-tar. The production of petroleum on the Caspian is only in its infancy.

Similar results have not yet been actually obtained from the residues of American petroleum, but it is highly probable that they also will prove available sources of aromatic compounds. Hence should even the last gas-works in the world come to a close there is no reason to fear a deficiency in the supply of benzol and anthracen.—*Die Chemische Industrie*.

ON SOME WATER FROM THE RIVER DART WHICH DESTROYED FISH.

By Dr. T. L. PHIPSON, F.C.S., &c.

In my notes of some analyses of waters recently published in the *CHEMICAL NEWS* I have alluded to the importance of the indication furnished by *phosphoric acid* in the analysis of potable waters: I stated that, having tested a considerable number of spring and river waters for this substance, I had found it generally absent, and that when present it is undoubtedly a bad sign. When I wrote that Paper I had not access to the whole of my Laboratory memoranda, and I have since found in my notes of 1875-6 an examination of a specimen of water from the River Dart, taken at a portion of its course where it was found to be injurious to fish, and killed large numbers of trout.

Instead of yielding some 15 to 22 grains of total residue per gallon, as we might expect in an ordinary specimen of river water flowing through populous districts, it gave a very much worse result. I found this water *somewhat turbid*, and it showed a *slight* deposit of organic matter (vegetable *débris*, fragments of *Algæ*); it was *neutral* to test-paper. It gave—

Total residue, 52.50 grains per gallon imperial.

	Grains.
Nitrogenous organic matter, burning with a bad odour; nitric acid, and ammonia ..	17.50
Phosphoric acid	3.25
Mineral matters: carbonate of lime (chiefly silica, sulphate of lime, alkaline salts, &c....	31.75

The strata through which this water flows is the chalk of Kent. The presence of so much organic matter and that of phosphoric acid to so large an amount points either to sewage contamination or to refuse from chemical works finding its way into the river near the spot where this sample of water was taken. The former suggestion appears the most probable. This analysis was made in December, 1875; and in February, 1876, I examined a large trout that had died in this water. No poison could be detected in the stomach and intestines; strychnine and picrotoxin were more particularly sought for. The gas contained in the air-bladder, which was much distended,

had a very foul odour (it is usually odourless, or nearly so); and at the commencement of the *oesophagus* I found a piece of printed paper, half swallowed and partially digested, bearing the date 24th November, 1875.

NOTICES OF BOOKS.

Elementary Lessons on Sound. By Dr. W. H. STONE, Lecturer on Physics at St. Thomas's Hospital. With Illustrations. London: Macmillan and Co. 1879.

THE speciality of Dr. Stone's well-written Manual is the large amount of attention devoted to the consideration of the connection between acoustics and music; in fact there is no elementary work that we know of in which this branch of the subject is so well and fully treated. Beginning with the different methods by which sound is produced—whether by strings, rods, plates, bells, membranous reeds, columns of air, heat, or electricity—it gradually leads up to the modes by which it is propagated, its velocity, wave-motion, reflection, and refraction. Intensity, consonance, and interference are well described. The chapter on Pitch is also a good one, the various methods of tonometry being very clearly given. The mixed question of standard pitch is also lucidly described, as well as the courses which have led to the gradual rise in pitch that has taken place in European orchestras.

Chapter V. is devoted to the consideration of the nature of musical sounds as distinguished from mere noises; and Chapter VI. to the effects of heat, atmospheric pressure, moisture, and density on sound in general. The difficult questions of scales, chords, temperaments, and tuning are clearly treated. We see no mention in this chapter of the Oriental scale, which differs from ours, and cannot therefore be noted according to our notation, although Eastern melodies can be reproduced on a perfect instrument, such as the violin. It may be briefly mentioned that, owing to the intervals between all the notes except the first and second being different to ours, harmony is impossible except in octaves. The different attempts of Peronnet Thompson, Pole, Bosanquet, and others to produce instruments with perfect temperaments are fully described and illustrated, as well as Mr. H. Bassett's ingenious comma and telephonic trumpets, in which perfect intonation is very closely approached.

An excellent description of the different kinds of musical instruments, and the principles upon which they act, is given in Chapter VIII. In this chapter there is a very clear and succinct description of the mechanism of the human ear, which the author very properly treats as a musical instrument. The same remarks apply to the description of the organs of the human voice.

The illustrations, of which there are nearly seventy, are generally good, but many of them seem to be badly over-printed, and some are from worn-out or damaged blocks. When will our English scientific publishers take a leaf out of the books of our American brethren in this respect? They are completely distancing us in the art of wood engraving as applied to scientific illustration, as any one may see by opening the "*Scientific American*," or any similar paper published on the other side of the Atlantic.

We should have been glad to see a fuller description of the telephone, microphone, and their congeners. In a work on Sound published in 1879 they surely deserved a better fate than being dismissed with exactly nine lines of mention. Prof. Tyndall's researches on the reflection of sound from layers of different temperatures also deserved fuller treatment. The phonograph, too, is dismissed in five lines, and Captain Galton's is not even mentioned, at least as far as we could discover.

With the exception of the woodcuts the book is well and clearly printed, and the frequent side-headings will be a great convenience. Beyond the few defects we have

pointed out we have no reason to say anything but in praise of Dr. Stone's little book, except that the index is a miracle of meagreness, and ought to bring down on his head the major anathema of the Index Society. Nevertheless we heartily recommend his book not merely to the science student, but to the musician who really wishes to be instructed in the scientific principles of the glorious art he practises.

CORRESPONDENCE.

TESTING COMMERCIAL POTASH.

To the Editor of the Chemical News.

SIR,—Some information recently gathered personally in the United States and Canada as to the commercial potashes produced there will, I think, be of interest to many of your readers. Being the managing director of a concern producing caustic potash free from soda, the subject naturally was one of great interest to me.

I visited the Montreal potash warehouses—a special warehouse where all the potashes produced in Canada are taken to be officially examined, tested, and branded, according to their quality as “firsts,” “seconds,” or “thirds.” I was told that the system of examination and testing employed was most complete, and that the official branding here of quality was universally accepted without question. The inspector informed me that he first inspected the general condition of the casks; that they were then opened and the contents examined for colour and appearance, and a carefully drawn average sample taken to ascertain the amount of potash they contained; that this was most exactly ascertained and calculated by a standard solution of sulphuric acid and litmus paper! On gently hinting to the official inspector that this was hardly a satisfactory manner of procedure, and that the only accurate way would be by also ascertaining the actual potash contents in the usual way by the platinum method, I was astonished to find that he was quite ignorant of the difference between the two alkalies, potash and soda!

It will be manifest to all your readers that there is nothing to hinder adulteration with soda-ash to almost any extent, and that that the potash will pass the inspection perfectly—a good 58 per cent soda-ash fused with the potash would easily come out as a “potash” of 78 per cent. Soda-ash is now worth in Montreal about £8 per ton, potash about £20 per ton, so that this source of profit will doubtless be availed of by enterprising producers of wood-ashes up in the country.

In New York and Philadelphia there is no official inspection, and out of the many consumers of wood-ashes that I saw and talked with in these districts, I only met one who actually tested his ashes for potash contents. There was a general complaint, however, that “somehow” their production was often unaccountably short or unsatisfactory, even though the ashes contained a full strength of potash according to the alkalimeter. The reason I think of this is very evident.—I am, &c.,

W. J. MENZIES,

Managing Director Greenbank Alkali Works Co.

St. Helens, Lancashire, July 23, 1879.

CHRONOLOGY OF THE ISOMERIC PURPURINS.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxix., p. 255, I find an article on the above subject in supplement to which I beg to say the following.

In December, 1871, Dr. Gessert of Elberfeld forwarded to Prof. E. Kopp, of the Technical Laboratory at Zürich, a bottle, labelled “artificial purpurin,” and requesting thorough researches as to the nature of the “purpurins” contained in the substance. Dr. E. Kopp entrusted me with the working out of the subject in question. The consequent investigations led to the method described by me in the *Moniteur Scientifique*, and it is in the same paper that the name of isopurpurin was given to the product separated from the original paste by my process. The investigations were published in March, 1872, and the results forwarded to Dr. Gessert, Prof. E. Kopp promising me at the same time to fully report the matter in the *Moniteur Scientifique*. I am unable to say why the publication was delayed from March to August, but it is highly probable that technical reasons were the cause.

Mr. Perkin's first preliminary publication of 1870 had not come to my knowledge at that time, and when the second preliminary report of 1872 appeared, my paper was already concluded. The full description of anthrapurpurin, however, only was brought out in 1873, fully a year after my paper on isopurpurin.

Regarding the pure state of iso- and anthra-purpurin at the time of their description (1872 and 1873), later researches in both the subjects have shown that neither of the bodies described in the original reports (1872, iso-, and 1873, anthra-purpurin) were the chemically pure ones. As to the isopurpurin, Prof. Morton has taken trouble in fully showing its having been a mixture in 1872, and, in fact, only in 1876 Messrs. Schunck and Roemer succeeded in obtaining this body in a pure state. The same chemists, however, find that the original anthrapurpurin of Perkin was by no means a pure body, and came to the conclusion that just the fact of its having been impure led to the error of its being “sparingly soluble in alcohol,” the very quality forming the main difference between the iso- and anthra-purpurin. Messrs. Schunck and Roemer found the pure anthra-purpurin “easily soluble in boiling alcohol,” (*Ber. der Deut.*, 1877, p. 680) and a foot-note says:—“Mr. Perkin describes his anthra-purpurin as sparingly soluble in alcohol and we found the same with a sample received from this gentleman. Probably this depends upon small quantities of impurities contained in the anthra-purpurin, and for this view speak the results of the analysis, and the small ability for crystallisation of the product, which is almost amorphous.”

The above will easily show that the iso- and anthrapurpurin could not both claim the name of pure bodies at the same time of their original description, and that the investigation of both bodies had been made at same time, independently from each other.—I am, &c.,

J. AUERBACH.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 1, July 7, 1879.

Identity of *Bacillus Amylobacter* and of the Butyric Vibrio of M. Pasteur.—P. Van Tieghem.—The author contests the identity of these organisms, which had been asserted by Prazmowski, of Leipzig. The details of the paper are micro-botanical rather than chemical.

Evaporation of Water under the Influence of Solar Radiations which have Traversed Coloured Glasses.—A. Baudrimont.—It appears from the facts described that coloured glasses exercise a real influence upon the evaporation of water, and that the quantity vaporised varies with the nature of the colours. Green

and red are in general the colours least favourable to evaporation. Their relative activity fluctuated. Yellow and colourless glasses were the most favourable. In the experiments of 1859 blue and orange, which are complementary colours, were almost equal in action. The water of the basin of Arcachon presented the singularity that colourless glass was scarcely more active than green glass and was far surpassed by red glass. Red glass, which allows a slight evaporation of water is the one whose colour is extinguished by the smallest photometric thickness, whilst yellow and colourless glass let more light pass and produce the greatest evaporation. Green is often inferior to red in its action, although it transmits a greater quantity of light. The explanation of these phenomena is not readily found. Does heat alone undergo a kind of analysis under coloured glasses, as is the case with light? Is coloured light ultimately transformed into heat? The author has long been of opinion that heat is essentially distinct from light, being due to a movement of the molecules, and not of their constituent elements, as is the case with light; the vibrations producing it are larger and less numerous in an equal time, and are propagated with less speed.

Thermo-chemical Study of the Alkaline Sulphides.—P. Sabatier.—The author describes the heats of formation and of hydration of the anhydrous sulphide of sodium, of the hydrosulphide of sodium sulphide, and of the corresponding potassium compounds.

A New Metal Discovered by M. Telleff Dahl.—An account of the discovery and principal properties of Norwegium. (See CHEMICAL NEWS, vol. xl., p. 25).

Commercial Trimethylamin.—E. Duvalier and A. Buisine.—Commercial trimethylamin is not, as M. Vincent considers, a simple product, but a very complex mixture, containing only from 5 to 10 per cent of trimethylamin, about 50 per cent of dimethylamin, besides mono-methylamin, mono-propylamin, and mono-isobutylamin.

*Verhandlungen des Vereins zur Beförderung des
Gewerbfleißes.* June, 1879.

Studies on Tempering Glass.—Dr. Schott.—An elaborate account of all the methods proposed for tempering glass. The paper does not admit of useful abstraction and its reproduction is specially prohibited.

Report on Dephosphorising Steel by the Process of Thomas and Gilchrist, and Proceedings of the Iron and Steel Institute for May 7, 8, and 9, 1879.—The author considers that though the process has not advanced beyond the experimental stage it is still important to have shown that under certain conditions phosphorus can be eliminated in the Bessemer converter.

Aniline-black.—Dr. Häussermann.—Referring to Dr. Kayser's paper, the author remarks that for the last ten years anilines containing mere traces of toluidin have been exclusively used in the production of aniline-blacks.

Bulletin de la Société Chimique de Paris,
No. 12, June 15, 1879.

Preparation of Methyl-formic Ether and of Pure Methylic Alcohol.—C. Bardy and L. Bordet.—The authors prepare the former compound by placing in a flask formate of soda, dried at 130° to 140°, and powdered, to which is added at once a mixture of methylic alcohol and hydrochloric acid, the three substances being in equivalent proportions, with a slight excess of methylic alcohol. To the neck of the flask is fitted a worm surrounded with cold water, which is not renewed. Above the worm is a delivery-tube, which leads the vapours into a second worm kept carefully cooled. The flask is plunged in cold water, which is gently heated to a boil. When the water surrounding the first worm rises to 45° the

operation is complete. In this manner, from 2 kilos. sodium formate, the authors have obtained 1610 grms. pure formic ether, the theoretical yield being 1764. The ether is easily saponified by means of a solution of caustic soda at 30°, the exact standard of which is known, equivalent quantities being taken.

Pseudo-uric Acid.—E. Grimaux.—If uramile reacted upon urea with elimination of water and ammonia there would be formed a body of the composition of uric acid. Elimination of ammonia, however, alone took place, and the pseudo-uric acid of Baeyer was formed.

New Method of Formation of Glycocol by means of Nitracetic Ether.—M. de Forcrand.—It appears from the researches of Meyer and Stüber that the hydrobromic and hydriodic ethers of the fatty series if heated with silver nitrite give rise to isomeric nitric ethers, which on reduction yield alkalies. The author has applied the same reaction to the brom-hydric ether of glycolic acid.

Action of Ethylen upon Benzol in Presence of Aluminium Chloride.—M. Balsohn.—Among the substances formed are ethyl-benzin, diethyl-benzin, and triethyl-benzin.

Active Principle of Insect Powder.—G. Dal Siè.—This powder is obtained from species of Pyrethrum, of different origin. In 1876, Jousset de Bellesme extracted from *P. carneum* an alkaloid. In 1878, M. Rother discovered an acid, or rather a glucoside, possessing insecticide properties. The author has obtained from the ethereal extract a crystalline acid, and from the alcoholic extract a resinous matter, which in contact with dilute sulphuric acid splits up into sugar and another product.

Moniteur Scientifique, Quesnewille.

July, 1879.

Periodic Law of the Chemical Elements.—D. Mendeleeff.—From *Liebig's Annalen*, 1872.

Notices of Foreign Researches.—E. Noelling.—These notes consist entirely of extracts from the *Berichte der Deutsch. Chem. Gesellschaft*.

Medical Properties of the Salicylate of Soda.—A lengthy discussion on the value of the salicylates in the treatment of gout and rheumatism.

Studies on Ultramarine.—T. Morels.—The composition of natural ultramarine is by no means constant. The carbonate of lime found in it by Gmelin to the extent of 28 per cent belongs to the gangue in which the lazulite is embedded. Natural ultramarine does not resist alum and acids, but is decomposed by them with liberation of hydrogen sulphide often more readily than are artificial samples. There is consequently no means of distinguishing the natural from the factitious product. Dilute acids decompose artificial ultramarine with the evolution of sulphuretted hydrogen and sulphurous acid. The rose-coloured kind give off sulphurous acid alone. Concentrated sulphuric acid is without action. Hot concentrated solutions of caustic alkalies turn the colour from blue to grey. Saturated solutions of alum decompose ultramarine slowly in the cold, but more rapidly with the aid of heat. At about 200° it is converted by acids into a violet ultramarine, which finally becomes red. Ultramarine may be heated to redness without being decolourised, and with certain precautions it may even be incorporated in melted glass, but it begins to be affected at 120° and becomes less brilliant. The complete analysis of ultramarine is very laborious. The author gives here a process sufficient for the requirements of the manufacturer, and showing the silica, alumina, total sulphur, and soda. The sample is first decomposed by fuming nitric acid. No sulphuretted hydrogen escapes, and the total sulphur remains in the liquid as sulphuric acid. The liquid is evaporated to dryness, the residue is taken up in a little water acidulated with hydrochloric acid, evaporated a second time, again dissolved in acidulated water, and the silica remaining

undissolved is washed with cold water upon the filter. In the filtrate the sulphuric acid is precipitated with barium chloride, when the weight of the barium sulphate shows the total weight of combined sulphur in the ultramarine. Alumina is then precipitated by ammonia, and in its filtrate, after removing baryta by means of sulphuric acid, the soda is determined as sulphate. The properties required by consumers of ultramarine are a deep and brilliant colour, fineness, tinctorial power, and resistance to alum and to acids. All these points are determined by comparative trials with a standard specimen. Papers or textile materials tinted with aniline-blues on exposure to the sun are bleached often in a few hours. No other blue employed for this purpose fades so rapidly. Papers, &c., tinted with ultramarine or with cobalt, leave a blue ash on incineration. If ultramarine has been used the ash is decolourised by the application of dilute acids, whilst the shade of cobalt, on the contrary, is brightened by this treatment. Papers, &c., blued with prussian blue, yield on ignition an ash which gives the reactions of iron.

On Uralium, a New Metal of the Platinum Group. A. Guyard.—As far back as 1869 the author discovered this metal in commercial platinum obtained from Russian ores. Next to silver it is the whitest metal known; its malleability is as great as that of the purest platinum, but its ductility is much greater, and it is almost as soft as lead. Its melting-point lies near to that of platinum, and it is not volatile. Its sp. gr. = 20.25, and its molecular volume, like those of osmium, platinum, and palladium, is 6.25. Its atomic weight has been found 187.25. In its chemical properties it is difficult to distinguish from platinum.

Flour mixed with Mineral Substances.—C. Cailletet.—The author's method for detecting the tenth of a milligram of alum, magnesia, chalk, gypsum, arsenious acid, &c., added to 10 grms. of flour, depends on the insolubility of the flour of wheat, rye, barley, &c., in chloroform; on their specific gravity, which is less than that of chloroform, and on the specific gravity of the mineral matters, which exceeds that of chloroform. He takes a perfectly dry glass tube 20 centimetres in height, and 2 to 3 in diameter. 10 grms. of the flour are introduced, the tube is nearly filled with chloroform, corked, and shaken for a minute. It is then let stand in an upright position, and in a cool place for some time. The flour which floats on the surface is removed, the chloroform is decanted off and may serve for new operations, and the deposit is treated with cold distilled water, which dissolves alum. The substances insoluble in water are collected on a filter, dried, weighed, and examined physically and chemically. Mineral salts existing naturally in the flour are not deposited, but remain in the floating layer.

M. Chevreul, now in his 93rd year, began his usual course of lectures on organic chemistry at the Museum of Natural History on June 10th.

Chemiker Zeitung.
No. 28, July 10, 1879.

According to Dr. F. Brockhoff a solution of magnesium chloride is found preferable to water for filling gas-metres. There is no appreciable loss by evaporation, freezing is practically impossible, and the gas is freed from ammonia.

The stearin manufacturers of Paris and its district, who represent one-third of the production of candles in France and employ 6000 men, have petitioned for the abolition of the duty upon candles.

Antiseptic Action of Acids.—According to Sieber a relatively small proportion of acid, 0.5 per cent, prevents putrefaction. This property is conspicuous in the mineral acid, and in acetic acid. Lactic and boric acids are much less effective.—*Journ. Prakt. Chemie.*

Chinese Galls.—A new kind of Chinese galls have been lately introduced. They are of the size of a plum, bent over at the point, and contain 72 per cent of tannin.—*Arch. Pharm.*, ii., 724.

Benzol and Benzin.—These names have generally been regarded as synonymous, but certain pharmaceutical works now apply the name benzin to a light petroleum product. True benzol is soluble in half to three-quarters its weight of alcohol, while the petroleum spirit requires six times its weight.—*Arch. Pharm.*, ii., 519.

Acid Reaction of Flowers.—It is not correct to assert that the juices of red flowers have always an acid reaction, whilst those of blue flowers react neutral or faintly alkaline. Vogel has shown that many blue flowers have an acid reaction. The red flowers of *Pisum sativum* are neutral.—*Akad. Wissen. Munich.*

No. 29, July 17, 1879.

The "new burette" constructed by Dr. Lagrange seems according to the description to differ little from that of Binks. It is bifurcate above, the wider aperture serving for the introduction of the solution, which is then dropped from the narrower orifice.

Revue Universelle des Mines, de la Metallurgie, &c.
Tome 5, No. 2, March and April, 1879.

This issue contains no chemical matter save what is taken from the *Comptes Rendus*.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 11, July 10, 1879.

Falsified Autographs.—The chief difficulty in the way of the ingenious forgers of such documents lies in the ink, which cannot be brought to exhibit the irregular fading of genuine ancient writings.

Curious Behaviour of Lightning.—A house in the Avenue de Clichy was struck by lightning, which followed a water-pipe to the earth and then re-ascended to the fourth story.

No. 12, July 17, 1879.

This issue contains no original chemical matter. There is a curious account of the visit of M. Moigno to Rome for the purpose of obtaining the papal benediction upon his "Splendeurs de la Foi."

Reimann's Fürber Zeitung,
No. 27, 1879.

The Rhenish papers announce the death of a girl from blood-poisoning, due to the alleged action of dyed stockings. Dr. Reimann points out that this case, like others of the class, is strictly anonymous, and without any guarantee of its authenticity.

La Correspondance Scientifique.
July 15, 1879.

Speed of Electric Currents.—According to the recent investigations of M. Siemens, described in the *Industrie Blätter*, the speed of an electric current, under the circumstances of the experiment, is 30.2 geographical miles per second, or about 40 kilometres.

Biedermann's Central-blatt.
June, 1879.

The Reverted Phosphoric Acid in Superphosphates.—Prof. A. Millot.—It is supposed that those phosphates are most important for the nourishment of plants which are soluble in water containing carbonic acid. The attempt has been made to determine their value according to their solubility in the citrate and the

oxalate of ammonia. The action of these reagents is disorientant. Certain aluminous and ferriferous phosphates dissolve very readily in the citrate of ammonia, but sparingly in the oxalate, whilst other phosphates behave in the inverse manner. These tests would lead us to assign a higher value to the phosphates of iron and alumina than to dibasic phosphate of lime. To determine the value of a superphosphate the author finds, first, the proportion of phosphoric acid soluble in water; secondly, the phosphoric acid soluble in acetic acid (from which that soluble in water is deducted), representing the bibasic phosphate of lime; lastly, determination of the phosphoric acid soluble in a cold ammoniacal solution of citrate of ammonia, from which again the two former quantities are subtracted. The author finds that the presence of ammoniacal salts prevents to a great extent the reversion of soluble phosphates.

Absorption of Mineral Matter by Plants.—P. Dehérain.—Plants whose roots are placed in a solution of sodium chloride, either unmixed or accompanied by small quantities only of potassium and calcium chlorides, absorb sodium in smaller proportion the more of the other salts is present.

On Schizomycetic Fermentation.—Dr. A. Fitz.—In the warm indigo-vat the indigotin is reduced by hydrogen liberated by fermentation. No trustworthy statements are to be found concerning the nature of the organism which excites the fermentation, but it is probably *Bacillus subtilis*. The liquid is first heated to a boil, then cooled down to 104° F., and kept at that temperature, exactly as if the propagation of *Bacillus subtilis* was intended.

Action of the Milky Juice of Carica Papaya.—Dr. L. Wittmak.—The author proves experimentally that the juice of this tree is (or contains) a ferment that acts energetically upon nitrogenous bodies, and, like pepsin, coagulates milk. Its action is more rapid than that of pepsin, and is not arrested by temperatures of 60° to 65°. If boiled it yields a precipitate, as also, on the addition of mercuric chloride, iodine and the stronger mineral acids. —*Naturforscher*.

Justus Liebig's Annalen der Chemie,
Band 197, Heft 3.

Communications from the Chemical Laboratory at Griefswald.—These consist of a memoir by Dr. L. Spiegelberg, on Nitro-, Amido-, and Bromo-sulphobenzolic Acids, with a description of many of their Salts; and a memoir by Dr. Heinzelmann and Dr. Spiegelberg, on Pentabrom-sulphobenzolic Acid.

Action of Dehydrating Agents upon Camphoric Acid and its Amides.—M. Ballo.—The author's object was to obtain the nitrile of camphoric acid, which agrees in its empirical composition with nicotin. He obtained a small quantity of a nitrile, $C_{10}H_{14}N_2$, a colourless crystalline body which in a state of absolute purity would probably be inodorous. It is insoluble in water, soluble in ether and alcohol, and sublimes between 125° and 130° without melting. It is apparently saponified on boiling with potassa.

MISCELLANEOUS.

The Rare Metals.—Dr. Theodor Schuchardt, of Goerlitz, sends us the following quotations for the rare metals to which we directed attention in our issue of the 18th ult.:—Cerium, 20s. per gramme; lanthanum, 40s. per gramme; and didymium, 30s. per gramme. These are obtained in globules by electrolysis. Thorium, in powder, is 36s. per gramme.

A Visit to Mr. Edison's Laboratory and Works.—Mr. W. Lant Carpenter has kindly sent us a copy of a letter to the *Bristol Mercury*, in which he gives an account of a visit to Mr. Edison, at Menlo Park. The following is an abstract of the letter:—The laboratory, workshops, &c., as well as some isolated buildings, for delicate electrical measurements, are spread over an acre of ground, railed in, admission to which is only given to privileged visitors. The machinery is driven by a beautiful, high-pressure, 80-horse engine, also used to drive the electric light machinery, most of which is in the same shop. About a dozen workmen were engaged, some in electrical test measurements, &c., but chiefly in manufacturing Mr. Edison's latest form of telephone, constructed for the electric and hygrometric conditions of our English atmosphere. About 50 of these have already been sent to the Edison Telephone Company, No. 6, Lombard Street, London. Their price in New York is 14 dollars per pair, or about £2 15s. The same instrument contains a mouth-piece for the transmitter, and a vibrating disc as a receiver, about 3½ inches diameter, from which the sound comes out loud enough to be heard all over the room. In this instrument the carbon button of the old Edison telephone is replaced by a cylinder of strongly compressed chalk, rendered a conductor by being moistened with a solution of, it is believed, phosphate of soda. A weak battery current is used in this, as in the other Edison telephones, the intensity of which is varied by the voice acting in some unexplained way upon the chalk conductor. Mr. Edison admitted that he was not doing very much at present at the problem of domestic electric lighting. He appeared to consider the question of its economical sub-division a solved problem (he had 16 lamps in the workshop, each with its small coil of platinum wire, in a glass globe, 3 or 4 inches diameter), and was now giving attention to the details of lamp construction. This new form of lamp is to be a minute cylinder of compressed pure zircon, which is to be heated to whiteness by the surrounding coil of platino-iridium wire. A chemist was engaged in purifying zircon for this purpose. There were only two electric light machines in the shop—one a discarded American instrument, the other Mr. Edison's own invention, apparently very simple, but giving a current in one direction only, not the rapidly alternating currents given by most machines. The newest thing in the shop was a dynamometer, the last and best of several invented by Mr. Edison, with the result of which he was perfectly satisfied, and he stated that with this instrument he had been able to show that, after deducting the necessary amount for friction in the machinery, more than 95 per cent of the mechanical force employed was obtained in the form of light. Time did not permit us to go through all the steps of the demonstration, which he ended by saying, "and, therefore, this is the most perfect machine in existence." As far as could be seen there appeared to be some fallacy (1) in the expression of mechanical force in terms of electricity, or *vice versa*; or (2) in the expression of light-intensity in terms of electricity, or *vice versa*. The principle adopted in the dynamometer is that of the sustaining motion which keeps a weight-driven clock going while it is being wound up. Upon the belt which drives the machine is placed a loose pulley, to which is hung a box, containing 1000 lbs. of iron. The belt is thus pulled down in the centre. The box is placed upon a weighing machine, and in normal conditions weighs 1000 lbs. When an additional resistance is put on, as by working an electric light machine, the weight of this box is altered, and from the amount of this alteration, and the known speed of the belt, the desired results are calculated. Mr. Carpenter also informs us that in consequence of enquiries by circular addressed to miners in Colorado (whence he has just returned) and elsewhere by Mr. Edison, platinum is found to be much more extensively distributed out there than was supposed.

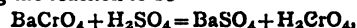
THE CHEMICAL NEWS.

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GREEN PIGMENT FROM CHROMATE OF BARIUM.

By THOS. DOUGLAS.

Barium chromate, which precipitates on the addition of a solution of barium chloride to a solution of a soluble chromate, is used to some extent as a pigment under the name of "lemon-yellow." When strong sulphuric acid is added to this substance in the dry state great heat is developed, and it is coloured a deep red from the liberation of chromic acid. If the mixture be now ground up in a mortar and heated to bright redness, the chromic acid is decomposed into the sesquioxide, which colours the mass green, yielding a pigment possessing considerable body. Assuming the reaction to be—



the amount of sulphuric acid requisite for the complete decomposition of barium chromate is 38.7 per cent. It is not advisable, however, to decompose the chromate completely; this would dilute the pigment too much with barium sulphate; and, besides, the presence of a certain proportion of the yellow element seems to improve the quality of the colour. Very good results are obtained by using 20 per cent of sulphuric acid (o.v.)

From the great heat developed it might be rather dangerous to mix the dried chromate with strong sulphuric acid on a large scale. The better method of proceeding would probably be to mix the wet precipitate with acid of ordinary strength, and dry the mixture at a moderate temperature. In this way the chromic acid would be thoroughly incorporated with the other substances.

The pigment produced by the above process would doubtless possess great permanence and freedom from the objections arsenical greens are liable to. The cost of production ought to be moderate.

Manure Works, Blaydon-on-Tyne.

TRANSFERRING LIGHTFOOT-BLACK FROM ONE FIBRE TO ANOTHER.

By JUSTUS WOLFF.

LIGHTFOOT-BLACK dissolves in a strong solution of aniline hydrochloride in water, but not completely, with a deep greenish black colouration. The solution obtained in that way added to hot water dissolves with black violet colour, and this liquid dyes cotton, wool, and silk grey. Even the Lightfoot-black on the fibre dissolves in a strong solution of aniline hydrochloride (produced by mixing 12 parts of hydrochloric acid of commerce and 10 parts of aniline).

About three years ago I dyed a large quantity of China-grass yarn with Lightfoot-black by soaking the yarn thoroughly in a strong solution of aniline hydrochloride and potassium chlorate, greening for about a week, and then passing through a bath containing small quantities of chrome and hydrochloric acid, washing, and drying. A small quantity of that yarn treated lately with a strong solution of aniline hydrochloride produced a dark greenish black solution, whilst the remaining fibre, after washing and drying, showed a dark greenish grey colour. (Of course the fibre was corroded by the action of the aniline salt. The greenish black solution mixed with water dyed cotton a beautiful bluish grey, and wool and silk a blackish

grey, all standing soap very well, that on cotton even improving by treatment with soap. We see here that this colouring matter has by itself a very great affinity for the fibres, without being produced on the fibre as in the Lightfoot process.

The shades thus produced on wool and silk are not bright, proving that the Lightfoot black process is unable to produce fine black shades at all on these animal fibres. The solutions obtained in the above manner contain too much acid and comparatively small quantities of colouring matter, so that it is very difficult to dye a deep black with. As far as I know, this is the first case of transferring Lightfoot-black from one fibre to another.

If the solution of Lightfoot-black in aniline salt solution is neutralised with caustic soda, diluted with water, and boiled till all the aniline is driven off by the water vapours, a greyish black powder remains in a light brown coloured slightly alkaline liquid. The powder filtered from the liquid, and washed on the filter with boiling water, consists of two different colouring matters, the one dissolving in boiling water acidulated with hydrochloric acid with a bright red colour, dyeing cotton and wool of a dull red shade, which by washing with clear water turns reddish brown, and by soaping clear brown, and the other consisting of a dark blue-black powder, insoluble in neutral and acidulated water.

Here we have another proof that the Lightfoot-black consists of two colouring matters—a dark blue and a brown one.

Wigan, July 30, 1879.

ON THE
EXPLOSION OF THE FLOURING MILLS AT
MINNEAPOLIS, MINNESOTA, MAY 2, 1878,
AND THE CAUSES OF THE SAME.

By S. F. PECKHAM.

As I was sitting at the tea-table on the evening of May 2, I was startled by a noise that sounded as if something as heavy as a barrel of flour had been tipped over on the floor above. A few seconds later the sound was repeated, and we all ran to the door which commanded a full view of the falls and manufacturing portion of the city. An immense volume of black smoke enveloped the spot where the Washburn A Mill had stood, and a perpendicular column of smoke was projected into the air above the elevator at least four hundred feet. The Humboldt and Diamond Mills were directly behind the elevator from the place where I stood. A heavy wind was blowing from a point a little to the east of north, a direction from the Washburn A Mill towards the elevator and the other two mills. In less than two minutes from the time of the first explosion, the elevator, which was 108 feet high, was wrapped in flames from top to bottom. If the structure had been saturated in oil the flames could not have spread much more rapidly. In five minutes flame and smoke were pouring from every window in the Day and Rollins, Zenith and Galaxy Mills, which were between the Washburn A Mill and the river, producing a conflagration which from ordinary causes would not have gained such headway in two hours. Six flouring mills, the elevator, a machine shop, blacksmith's shop and planing mill, with a number of empty and loaded cars, were in flames in five minutes from the time fire was first observed by any one who survived the disaster.

From my own point of observation, which was about a mile distant, but two distinct explosions were heard; others nearer heard three, the first not as violent as the other two; while those nearer still heard in addition a sound which they described as a succession of sharp hisses, resembling the sound of burning gunpowder.

Those observers to the windward whose attention was arrested by the light produced, beyond the distance of half a mile, heard only one or two reports or failed to hear any report at all. From all the testimony in reference to sound it appears that the blow upon the air was not sufficiently sudden to produce a penetrating sound, but rather a dull, heavy blow, which was not communicated laterally to any great distance.

Burning wheat or flour was smelled for several minutes before the explosion by persons in such a position that the wind would carry the odour to them. Smoke was also seen issuing from what was known as the exhaust flour-dust spout of the Washburn A Mill for several minutes preceding the explosion.

At the instant the explosion occurred all observers agreed that the Washburn A Mill was brilliantly illuminated from basement to attic. The illumination was reflected from the water at and around the falls in such a manner as to remind one observer of the effect of a brilliant sunset. Another compared it to the reflection of sunlight from windows when the sun is near the horizon. Still another, who was crossing the lower bridge, had his attention called to what appeared to be a stream of fire, which as he described it, issued from a basement window and went back again. Immediately thereafter each floor above the basement became brilliantly illuminated, the light appearing simultaneously at all the windows, only an appreciable interval of time intervening as the stories ignited one after the other. Then the windows burst out, the walls cracked between the windows and fell, and the roof was projected into the air, followed by an immense volume of smoke and flame which ascended to an estimated height of from six to eight hundred feet. As the column of smoke was expanded and borne off upon the wind, brilliant flashes resembling lightning passed to and fro.

Two men, so near the Humboldt Mill that they were nearly buried by the falling rubbish, and on the opposite side from the Washburn A Mill, heard a loud report distinctly while the walls of the Humboldt Mill were still standing and at the same time were knocked down. Immediately after they saw flames issuing from the basement windows of the Humboldt Mill and at the same instant, before they could regain their feet, they experienced a second shock and miraculously escaped being buried beneath the falling walls.

The enormous and sudden displacement of air which followed the explosion, and the tremendous force which was consequently exerted laterally, was shown in the condition of the round-house of the Chicago, Milwaukee, and St. Paul railroad, and the broken windows in all directions. The round-house was a wooden structure about forty or fifty feet from the Diamond Mill. The sills were drawn out toward that mill until the building burst, letting a part of the roof fall in and leaving the sides standing at a sharp angle. Ordinary windows, and those of strong plate-glass on Washington avenue, one-fourth of a mile distant, were projected into the street. Not only the glass but the sash went out bodily, particularly in the lower stories of the buildings. Persons on the river at the water's edge noticed a displacement of the water producing a wave estimated to be eighteen inches high, before they heard the report of the explosion.

Whole sheets of the corrugated iron with which the elevator was covered, measuring eight by two feet, but quite thin, were picked up on the east side of the river more than two miles distant, and pieces of six-inch flooring from two to ten feet long were carried to intermediate points.

An examination of the ruins of the several buildings showed that the walls of the Humboldt Mill lay upon those of the Diamond Mill, and those of the Diamond Mill upon those of the west end of the Washburn A Mill, *showing that the buildings did not explode simultaneously but successively. The Washburn A Mill evidently exploded first from fire originating within it, and the high wind pre-*

vailing at the time carried the *flame* into the adjoining mills to the south, and away from the mills next the river. There was enough burning middlings and flour thrown through the broken windows of the latter mills to set them on fire, but they did not explode. Some significance may attach to the fact that the three mills that exploded were all running with more or less open French middlings purifiers, while the three that did not explode had been shut down for several days. There is no question but that the French purifiers project a great deal more dust into the atmosphere of the mills than those that are enclosed, but I have no doubt that in *any* flouring mill sufficient dust accumulates upon beams and machinery to produce an explosive atmosphere if from any cause this dust is scattered into the air and flame is communicated to the mixture while the dust is suspended.

There was less than a barrel each of lard oil, lubricating oil, and high-test kerosene in the Washburn A Mill at the time of the explosion.

There is absolutely no proof that any explosive material other than is produced in the manufacture of flour from wheat was in any one of the buildings destroyed, in the cars around them, or in the neighbourhood. The testimony of millwrights conclusively showed that fire produced by heated bearings is of such extremely rare occurrence in flouring mills as to practically exclude such a cause.* No suspicion of incendiarism has ever been expressed.

A slight fire, the effects of which were in no wise serious, occurred in the Washburn A Mill about three months before the explosion. It was discovered from the outside of the mill that smoke was issuing from a spout or conductor that discharged the air that was drawn through between the stones. The object for which the air is drawn through is to cool the stones and to carry off the vapour produced from the wheat by the rise of temperature due to friction. In this case the effects of *fire* were traced back from the outside of the building to one of the sets of stones on the north side of the mill used for grinding middlings. The effects of *flame*, however, did not extend beyond the blower which produced the exhaust. This led to the conclusion that the fire did not enter the dust-house, although the smoke must have passed through it. It is supposed that the fire was caused by friction between the stones, they having run dry from one of the causes that may produce dry stones.

In answer to enquiries made of several millers in the Minneapolis Mills, I found them uniformly of the opinion that the meal or flour as it left the stones had a temperature of about 100° F. or less. A number of careful experiments, made with an ordinary chemical thermometer, showed that the wheat enters the stones from the dryers at a temperature of fully 100° F., and that it leaves the stones at 120°—130° F. The temperature of the ground middlings as it left the stones averaged about ten degrees higher.

It was also the concurrent testimony of millers and mill owners that dry stones are of comparatively frequent occurrence, and that they are practically unavoidable. I am convinced that in the Washburn A Mill the frequency of danger from dry stones was considerably increased in consequence of the large number of stones in the mill, and especially from the fact that so few men were employed having the immediate oversight of the stones. Only two men were employed at the same time for the forty-two run of stone, a number inadequate for that supervision which so important a matter demands, as it is impossible from the large space occupied by so many stones, and the noise incident to their action, that even with the usual signals employed dry stones should be detected as soon as they become a source of danger.

Obstruction of the feed from any one of a number of accidental causes will produce dry stones. The danger

* These gentlemen concurred in the statement that the spindle which carries the stone had been known to become *welded* into the socket in which it revolved, stopping the stone. When asked if the friction produced a welding heat, one replied, "No, nowhere near it." It must be an example of perfect metallic contact, producing cohesion.

arises from the friction of the stones heating the last portion of the grist that remains between the stones to a temperature sufficient to char it, or convert it into a substance resembling tinder, which would readily ignite from a spark produced by the stones striking together. Another source of danger arises from nails or gravel passing between the stones with the grist and increasing the friction, producing either a rise of temperature or a train of sparks; perhaps both.

I am aware that numerous instances of dry stones can be cited that have proved perfectly harmless. An instance is on record in which a run of stone ground each other all night with no other result than the complete removal of the grooves which gave the stones a cutting face. On the other hand, cases have occurred in which the grooves became filled with charred wheat of a dark brown colour, packed into them so solidly as to require a mill-pick for its removal. It requires no argument to show that this tinder thus formed would become ignited from a train of sparks that would inevitably follow contact of the stones as the grist became compacted or completely removed from between them. It was found by experiment* that masses of flour that had become heated and charred, ignited readily and smouldered, but were inflamed with considerable difficulty; but it should be borne in mind that a number of sets of these stones are connected with a common spout or conductor, through which a strong current of air is being continually drawn, and which is filled with a dense cloud of very fine particles of starch (chiefly) heated to a maximum temperature of 140° F. Experiment also proved that the proper mixture of flour dust and air would not burn explosively except when brought in contact with flame. White-hot wires and glowing charcoal only burned the particles in contact with them. But it was found that burning pellets of charred wheat and flour would ignite wood, which a strong draft of air readily fanned into a blaze. Under the conditions previously stated with a draft of air passing through the dry stones strong enough to convey the pellets of smouldering tinder into the common wooden conductor an explosion becomes possible.

It is urged that these conductors are damp from condensed moisture, and also that a large amount of moisture escapes from the wheat and is conveyed away by the current of air. This loss is, no doubt, correctly estimated at from five to six per cent. It is, however, chiefly during the first grinding of the raw wheat that this loss is experienced. The middlings is drier, is ground at a higher temperature and is ground finer, producing more dust. The higher temperature renders the material more inflammable, and at the same time ensures a more complete solution of the vapour in the current of air. Moreover, the first fire in the Washburn A Mill was traced directly to a set of stones which ground nothing but middlings, and all that is known concerning the origin of the fire that produced the explosion confirms the supposition that that fire originated in a set of stones on the opposite side of the mill, which was one of six sets, all of which were used exclusively for grinding middlings, discharging into a common spout or conductor which communicated directly with a dust-house in which the dust settled to the amount of several hundred pounds a day. An explosion in this conductor, communicating flame to the dust-house, would scarcely fail to cause the successive explosions of the dust-house and the different stories of the mill, the shock of the first explosion being sufficient to throw the dust of the mill into the air.

The opinion expressed by one of the witnesses at the inquest, "that stones are liable to run dry at any time by accident," and that "dry stones can hardly be avoided by any amount of foresight," appears to be generally entertained by millwrights, millers, and mill-owners. Let it be granted that all experience shows that ninety-nine per cent. of dry stones injures nothing but the stones

themselves, the one per cent. of residue is burdened with fearful possibilities. If dry stones cannot be prevented in small mills, where one miller has charge of perhaps six run of stone, the danger is more than proportionally increased in a mill where one man has charge of twenty run, both with reference to prevention and detection. The problem, therefore, for the consideration of parties immediately interested is how to prevent or detect dry stones, particularly those used for grinding middlings. This practical problem appears to be fundamental, and one compared with which all others are without much importance. It is true that but few millers are without their experience of minor explosions or flashes resulting from careless use of lanterns or open lights. Indeed, I have been profoundly impressed with the generally innocent reputation of flouring mills when considered in the light of the immense number of accidents well known to millers and insurance companies; a number surprisingly large if confined to those occurring in the States of Minnesota and Wisconsin within a few years past. The remedy in such cases is so obvious that the most ordinary care and intelligence is sufficient.—*American Journal of Science.*

University of Minnesota, Minneapolis.

UPON THE DETECTION AND ESTIMATION OF NITROUS ACIDS IN POTABLE WATERS, ACIDS, &c.

By ALBERT R. LEEDS.

II.—WITH POTASSIUM IODIDE.

THE well-known method of Trommsdorf is of a colorimetric character.* The potable water to be examined is treated with a solution of zinc iodide, starch, and sulphuric acid, which develop, in case nitrous acid is present, a blue colour. The amount of nitrous acid corresponding to this colour is found by striking in distilled water containing these reagents the same tint with a standard nitrous acid solution. Ordinarily, the unknown and known solutions are prepared at the same time and placed side by side, in order that they may be subjected, as nearly as may be, to the same conditions. It is essential, in fact, that they should be equally exposed to the action of the air, and should be acted upon by light of equal intensity and for an equal length of time. To point out the importance and necessity of these precautions is the object of the present paper. Our attention was drawn to them in the course of preceding studies, upon the rate of change of soluble iodides in the presence of free acids, and the results of those studies have been given at length (*Journ. Amer. Chem. Soc.*, vol. i., 1879).

Briefly to recapitulate, these results are as follows:—*In the dark*, access of air being freely permitted, the amounts of iodine set free, the amount of free acid present being the same in every case, increases with the increase in percentage of dissolved iodide. *In the light*, the iodide liberated, in the case of solutions of equal strength, increases with the amount of exposure to the light and the intensity of the illumination. When air is entirely excluded no decomposition occurs even at temperatures above the boiling-point, and when exposed for days to the direct light of the sun.

These conditions of change having been established, it became of interest to determine in what way the mimetic method of Trommsdorf could be replaced by an absolute mode of volumetric estimation.

As a preliminary determination, the limits of sensitiveness of the zinc-iodide starch solution were investigated, and the rapidity with which it would undergo change in presence of acid in diffused daylight. Nine comparison-

* Experiments made by Prof. L. W. Peck before the coroner's jury.

* *Zeit. f. Analyt. Chemie*, viii., 358.

tubes were placed in the comparator-frame, and in each was introduced 100 c.c. water, 1 c.c. H_2SO_4 (free from nitrous acid) and 3 c.c. zinc-iodide starch solution. This last solution was prepared by adding to 20 grms. ZnCl_2 and 5 grms. starch, 2 grms. ZnI_2 and 1000 c.c. water. The reagents were prepared in the laboratory, and were perfectly pure. To the nine tubes were added 0.2, 0.4, 0.8, 1.00, 1.50, 2.00, 2.50, 3.00, and 5.00 c.c. of a standard solution of potassium nitrite, containing per 1 c.c. 0.01 m.grm. N_2O_3 . In other words, the first tube contained 2 parts; the ninth, 50 parts of N_2O_3 in 100 million. This last changed immediately, the others as follows:—

With	3 c.c. N_2O_3 solution, after 10 minutes.
" 2.5 "	" 15 "
" 2.0 "	" 18 "
" 1.5 "	" 50 "
" 1.0 and 0.8 "	" 1 hour.
" 0.4 and 0.2 "	" 1 " 50 mins.

A tenth tube, containing 100 c.c. H_2O , 1 c.c. H_2SO_4 , and 3 c.c. ZnI_2 , and *no nitrite*, at the end of two hours had acquired a faint blue colour; in depth, however, inferior to that given by the 0.002 m.grm. N_2O_3 in the first tube. It will be seen from this table that the reaction cannot be applied to waters containing as much as 0.05 m.grm. N_2O_3 in 100 c.c. Perhaps 0.04 m.grm. in 100 c.c. is the superior limit, waters containing larger amounts than this requiring suitable dilution. And, moreover, since the very dilute solutions did not change at the expiration of intervals increasing progressively with the order of dilution, the determinations in these cases are affected with a corresponding degree of uncertainty. This is still more strikingly the case when ordinary potable waters, which are seldom colourless and contain various substances in solution, are used. For example, we have known the water of the Passaic River, which contained, according to a determination on one sample, by means of meta-diamido-benzol, 1.2 pts. N_2O_3 in 10 millions, to remain unchanged, on the addition of zinc-iodide starch and sulphuric acid, at the expiration of eight hours. After the same interval, distilled water, containing the same reagents, but no nitrite, had acquired a well-defined blue tint.

Apparatus for Absolute Volumetric Determination.

The preceding considerations, having demonstrated the importance of comparing the results obtained by the colorimetric method with those which would be gotten by some mode of absolute volumetric determination, an apparatus for effecting the latter object was constructed as follows:—Two comparison-tubes were fitted up in a manner similar to wash-bottles, except that the first comparison-tube was made to slip up and down upon its exit-tube. In this way its exit-tube might be forced down into the bottom of the first vessel when desired. The reagents were placed in the first comparison-tube; 100 c.c. of the liquid to be tested in the second. After a current of carbonic acid, washed by passage through a potassium iodide solution, had passed through both vessels for an hour, the exit-tube of the first vessel was pushed down, as above described, and the reagents forced over into the liquid under examination. In this way, the air dissolved in the reagents was swept out of them before they were allowed to come into contact with the unknown solution. A slow current of carbonic acid was maintained for two hours, at the end of which time the liquids were titrated. The reason for allowing so long an interval to elapse before titration was that it had been found previously to be necessary, in order to permit so small a quantity of N_2O_3 as 2 parts in 100 millions to decompose their entire equivalent of potassium iodide. So effectually does the exclusion of air prevent the liberation of iodide under the influence of light alone, that no disturbing influence is exerted upon the results, even when the comparison-tubes are exposed to the direct rays of the sun for two hours or longer.

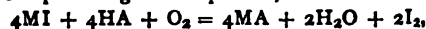
Example:—100 c.c. containing 0.05 m.grm. N_2O_3 were

introduced into the second comparison-tube, and 6 c.c. zinc-iodide starch and 1 c.c. H_2SO_4 in the first. After complete displacement of the air dissolved in reagents and solution, the reagents were forced over, and the current of carbonic acid gas continued. On titration with sodium hyposulphite, at the end of two hours, the amount required was 0.032 c.c. = 0.160 m.grm. I. The theoretical amount was 0.0334 c.c. = 0.167 m.grm. I = 0.05 m.grm. N_2O_3 .

Effect of Colouring Matters.

These experiments give the key to an explanation of the effect of colouring matters and organic impurities upon the rate of change of acid solutions of the soluble iodides. They suggest, likewise, a method of eliminating the perturbations so arising from titrations in which iodides are employed.

Colouring matters, &c., disturb the identity of conditions under which the colorimetric titration is made. Such an identity is more especially important in a determination like this, inasmuch as the liquids are undergoing a slow spontaneous alteration by contact with light and air. All colouring matters absorb more or less of the actinism of solar and other light, and, consequently, the amounts of iodine set free will differ correspondingly in solutions containing them. This absorbing action is especially great in the case of yellow solutions which, like caramel, cut off the more refrangible rays of the spectrum. If in addition to being coloured, these foreign bodies are decomposable and capable of combining with any of the oxygen of the dissolved air, their disturbing influence is two-fold. In the first place, they alter the rate at which acid solutions of soluble iodides decompose under the action of diffused or direct light, and secondly, they remove a portion of the oxygen which is essential to the progress of this decomposition. The conditions governing the reaction are summed up in the general equation,—



where M stands for any monovalent basic radical, and A a monobasic acid radical. (See "Influence of Light upon the Decomposition of Iodides," *Journal Amer. Chem. Soc.*, vol. i., part 3, by the author.)

Illustration: These suppositions were confirmed by the following experiment: Four comparison tubes were prepared, each containing 1 c.c. KI sol. (20 p. c.) + 1 c.c. H_2SO_4 + 100 c.c. H_2O + 0.05 m.grm. N_2O_3 . Enough caramel solution was added to each to communicate a strong colour. All four were exposed to sunlight for one hour, but through two of them a current of washed carbonic acid was passed, while the remaining two solutions were in direct contact with air. The results were as follows:

Caramel Solutions.

In Air.

- I. Required 0.95 c.c. $\text{Na}_2\text{S}_2\text{O}_3$ = 1.4 m.grm. N_2O_3 .
- II. " 0.90 " " = 1.35 " "

In Carbonic Acid.

- III. Required 0.05 c.c. $\text{Na}_2\text{S}_2\text{O}_3$ = 0.099 m.grm. N_2O_3 .
- IV. " 0.05 " " = 0.099 " "

It will be seen, that even where carbonic acid was employed, the amounts of N_2O_3 which were found are twice those of the A_2O_3 used. This excess was due to using reagents, out of which the dissolved air had not previously been driven by a current of carbonic acid. When this was done, by means of the apparatus before described, the results agreed with the theoretical. In case of the solutions exposed to air, the discrepancy is enormous, being nearly thirty times too great. Similar results were obtained in diffused light, but a greater length of time was required to produce them.

Conclusions.—The following conclusions are to be drawn from the preceding experiments:

- I. When the solutions to be titrated are colourless, and contain no compounds, organic or inorganic, which would

affect the percentage of dissolved oxygen, the determination may be made by the colorimetric method.

II. When the solutions are coloured, or contain organic or other bodies capable of absorbing oxygen, the air must be entirely removed from the liquids in order to obtain good results, and an absolute volumetric determination is essential.

With regard to the first point, it is to be remembered that, in any case, the colorimetric method less and less approximates to accuracy as we approach the lower limit of the reaction. This is due to the difficulty of securing equivalent amounts of change in excessively dilute solutions, when these solutions contain iodides in presence of free acid, and are not at the same time absolutely identical in tint and constitution. And, with reference to the second point, inasmuch as the colouring matter cannot always be removed from potable waters, by throwing them down through the formation of an insoluble precipitate, it is of utility to possess a method of determining nitrites in presence of colouring matters, and sometimes of certain classes of organic impurities.

Examination of Hydrochloric Acid for Chlorine, and of Nitric and Sulphuric Acids for Nitrous Acid.

After the preceding statements, it will be easily seen that the ordinary rule given in works on qualitative analysis, for the examination of sulphuric acid (*i.e.*, dilution with 20 pts. water and addition of potassium iodide and starch results), gives erroneous results when employed to detect minute quantities of nitrous acid, and the solution is allowed to stand for some time. But if the trial is made in an atmosphere of carbonic acid, care being used to expel previously all traces of air in reagents and in the dilute acid solution, the test and estimation of percentage may be satisfactorily performed. The same remark applies to the examination of hydrochloric acid for chlorine, by means of a soluble iodide. In the case of nitric acid, however, it is well not to expose the acid when under examination too long to the action of strong light, owing to the liberation of oxygen from the nitric acid itself under these circumstances. The ease and certainty with which the Griess reaction for nitrous action is performed should cause it to replace the former methods in most cases. So delicate is it, that the acids sold as chemically pure usually give a decided colour when the meta-diamidobenzol is added to them, and the general introduction of this reagent into use will probably raise the standard of so-called purity, at least in this respect.

Upon the Degree of Concentration at which Soluble Iodides will cease to remain unchanged in presence of Free Acid, even when out of Contact with Air.

The method pursued in examining this question, was to employ solutions of iodides and acid of gradually increasing concentration, until a point was attained at which decomposition ensued, even when all traces of air were rigorously excluded. The reagents were placed in the first, the iodides in the second comparison-tube, of the apparatus before described, and when all the air had been expelled from both, by the long-continued passage of carbonic acid, the reagents were forced over into contact with the iodides. In the first experiment, 1 c.c. KI (10 p. c.) + 5 c.c. H₂O and 1 c.c. H₂SO₄ (chemically equivalent to the 1 c.c. KI) + 5 c.c. H₂O, were employed. After expulsion of air, the solutions stood one hour in diffused light, and half-hour in sunlight, a slow current of carbonic acid flowing. On disconnecting and adding starch, no colouration took place.

In the second experiment similar solutions were used, but instead of being diluted to the 11th, they were diluted only to the 10th. They were exposed to diffused and direct sunlight for the same intervals. Before addition of starch, no change of colour was perceptible, but afterwards, a brownish-red. This compound was entirely

unlike in appearance to that formed by the union of free iodine with starch, under ordinary circumstances.

III. The reagent contained 5 c.c. KI (10 p.c.) + 2½ c.c. H₂O; in other words, they were now diluted only to one-tenth. After exposure for one-and-half hours to direct sunlight, they were brought into contact, when the same brownish-red precipitate was formed.

IV. No water was employed, but in its place 2½ c.c. starch-water, with the same amounts of sulphuric acid and potassium iodide. They were exposed, after expulsion of air, to diffused light for one hour, and then to sunlight for one-and-half hours. No change occurred in either comparison-tube until their contents were mixed, when the brownish-red precipitate formed immediately. On the addition of sodium-hyposulphite, the colour disappeared, showing that it probably contained either iodine or hydriodic acid. On collecting and washing the precipitate on a filter, it turned blue, and presented the ordinary appearance of iodide of starch.

V. A final experiment was made by dissolving 10 grms. KI in 30 c.c. starch-water. 20 c.c. of this solution were placed in the first comparison-tube, 10 c.c. concentrated H₂SO₄ in the second, and both treated with carbonic acid. The tubes were exposed to direct sunlight. In a few minutes the first comparison-tube exhibited a brown colouration, and after several hours a carmine-red precipitate was formed. After being allowed to stand over night, the red precipitate was still present, but another also, of a darker hue, lay above. On allowing the carbonic acid to flow through both precipitates for six hours, in the sun-light, they disappeared, and the original brown colouration only was visible. During this part of the experiment, the iodide and starch, it will be noted, were not mixed with the acid. Finally, they were driven over, when immediate decomposition ensued, and a large amount of iodide was set free.

Conclusions.—1st. In the absence of air and presence of carbonic acid, decomposition of an acid solution of potassium iodide occurs, when the concentration has attained to some point between one-third and one-tenth the weight of the water employed. 2nd. During the course of these experiments, a compound of starch has been formed which, from its deportment with reagents, and more especially from its turning blue on absorption of oxygen, may probably be regarded as a *hydriodide of starch*. An attempt would have been made to study it further had it not been for the difficulty of satisfactorily isolating a body which changed so readily on exposure to air into ordinary starch iodide.—*Journal of the American Chemical Society.*

CONTRIBUTIONS TO THE ESTABLISHMENT OF A RATIONAL SYSTEM OF EXAMINING WATER AS REGARDS ITS INFLUENCE UPON THE HEALTH OF MEN AND ANIMALS.

By Dr. F. HOLDEFLEISS.

(ABSTRACT.)

THE author, after a careful survey of the methods proposed and the results obtained, concludes that it is rarely practicable to determine the character of a water satisfactorily by chemical analysis alone. Microscopic examination, though too much neglected, promises more satisfactory results in as far as the organisms observed have been found to bear certain relations to the sanitary value of the water. The following points may be considered established:—

Only natural, freshly-drawn water, and the deposit taken from the bottom or sides of the well, tank, &c., should be submitted to examination.

Not the number of the organisms present, but their nature, must be regarded as decisive of the state of the water.

For the object in view the organisms present in water may be divided into three classes:—

(1.) Such as can live only in sound good water, which in any appreciable extent shows no suspicious decomposition of organic matter. Such organisms perish as soon as deleterious products of decomposition are met with in the water. Among this class are included the diatoms or Bacillaria, which, however, indicate a wholesome condition of the water only when found living. If we discover in a water merely the silicious shells of diatoms without their coloured contents, this is *per se* a certain sign of the suspicious character of the water. It is not necessary to determine the species, but merely to ascertain whether the shells contain protoplasm of normal texture and colour. *Spirogyra*, *Cladophora*, and other green thread-like Algae live only in good waters. They die off, their green chlorophyll shrinks up, and disappears when injurious decomposition products are introduced. They leave, however, no silicious shells behind, but perish altogether. The Desmidiæ behave in a similar manner. All these organisms, by means of their constituent chlorophyll, decompose the carbonic acid dissolved in the water for their nourishment.

(2.) Organisms free from chlorophyll, nourished solely from putrescent organic matter, and directly promoting the decomposition of organic matter, nitrogenous or non-nitrogenous. Such organisms can live only in waters which evince processes of putrefaction and an unwholesome, dangerous character. The presence of this group pre-supposes decomposing matter. Among these must be included *Beggiatoa alba*, a colourless alga, whose presence appears to be closely connected with the formation of hydrogen sulphide. *Beggiatoa* is found in bad well-waters, whilst the diatoms and other algae cannot live in covered wells, as they require light. Very similar is *Leptomitius lacteus*, which, however, gives off no sulphuretted hydrogen. We must also include *Crenothrix polyspora*, *Selenosporium*, and *Zooglaea*. Here likewise must be classed the Schizomycetes, the bacteria, monads, vibriones, &c., minute vibratory corpuscles which are developed in especial abundance where nitrogenous matter is plentiful. Hence the danger to be apprehended may be estimated by their abundance. The microscopic indications may here be usefully verified and supplemented by the highly sensitive tests for ammonia and nitrous acid.

(3.) In an intermediate group we place organisms which can live either in good or bad waters, such as the Oscillariæ, *Euglena viridis*, and the majority of the larger Infusoria.

We must distinguish between the water from (covered) wells and open waters, whether flowing or standing, and again between such as, up to the taking of the sample, have been entirely secluded from the light, and such as have been at least temporarily exposed to the sun, such as spring-waters.

Well-water is to be regarded as good when free from all organism, from ammonia, nitrous acid, and hydrogen sulphide.

Open waters, flowing or standing, are good when they contain living green algae and diatoms with the contents of the shells normally coloured, but no colourless algae (*Beggiatoa*, *Leptomitius*, &c.), and no, or but few, Schizomycetes.

Drinking-water for human use, in addition to freedom from all organisms, from ammonia, nitrous acid, and sulphuretted hydrogen, should not exceed 18–20° of hardness. If river-water is so purified by filtration that it possesses these properties, and that suspicious organisms do not reappear on standing, it may be used without scruple.

Open waters containing green algae and living diatoms and free from ammonia and nitrous acid may be consumed with safety.

For cattle traces of ammonia and nitrous acid in pond-

and river-water may be tolerated if green algae and living diatoms are present.

Water for fish-ponds should be free from sulphuretted hydrogen and should be rejected if *Beggiatoa alba* is present in a state of normal vitality. *Leptomitius lacteus* is probably dangerous.

The mere vicinity of waters containing *Beggiatoa* and the Schizomycetes is dangerous, as they may infect the atmosphere and communicate germs of a dangerous nature to the ground-water and the wells.—*Biedermann's Central-Blatt*.

NOTICES OF BOOKS.

Proceedings of the Twenty-sixth Meeting of the American Pharmaceutical Association, held at Atlanta, Georgia, November, 1878. Philadelphia: Sharrman and Co., 1879.

THIS is the first time that the American Pharmaceutical Association has held its meetings in the Southern States where pharmacy has hitherto been generally neglected. The present volume, containing nearly a thousand pages, is marked by the same thoroughly practical character that we have had occasion to praise in preceding notices. There does not seem to be a single paper or extract which does not contain numerous items of information calculated to help the working pharmacist in his everyday labours. During the past year, the Committee appointed to revise the U.S. Pharmacopœia appear to have done good work. The following recommendations have been made with regard to the new edition, viz., that the language used should be English, and that the present division into Materia Medica and Preparations should be abolished, and an alphabetical arrangement adopted. A list of the proposed elisions and additions is also published, and the comments of the medical and pharmaceutical professions are invited. Temperatures will be expressed both in F. and C., and double formulæ will be given in the case of chemical preparations of definite composition. Measures of capacity are to be entirely abolished and weights substituted. Other important additions are to be made in the form of tables. One of the most important reports was that read by Mr. C. L. Diehl, on "Fluid Extracts by Percolation," which is quite exhaustive in its character, a remark which applies equally well to that by Dr. Squibb, on "Fluid Extracts by Re-percolation." The Annual Report on the Progress of Pharmacy during the year ending June 30, 1878, extends over 644 pages, being more than double the length of that of last year. The next meeting will be held at Indianapolis.

CORRESPONDENCE.

ASHES OF WHEAT BRAN.

To the Editor of the Chemical News.

SIR,—In your issue of June 20 (CHEMICAL NEWS, vol. xxix., p. 276), just come to hand, I notice a criticism by J. Carter Bell, of the article "On the Ashes of Wheat Bran" which appeared in your number for June 6th, of which I am the author.

I beg to call Mr. Bell's attention to the fact that he has been analysing one thing while Miss Brown analysed another. By comparing notes, he would have informed himself that he has been examining "bran from white English wheat," while Miss Brown examined an ash residue of Minnesota wheat bran from under a boiler. No phosphate of alumina was given among the results of the analysis, because a careful examination of the substance analysed failed to reveal the presence of an appreciable amount of alumina.

The amount of sulphuric oxide (SO_3) should read 1.151 per cent., and the total 99.259 per cent.

While thanking Mr. Bell for his courtesy, I venture to suggest that in future, before he attempts the rôle of critic he ascertain what play is being played.—I am, &c.,

S. F. PECKHAM.

July 16, 1879.

EXPLOSIONS IN FLOUR MILLS.

To the Editor of the Chemical News.

SIR.—I have noticed in late numbers of CHEMICAL NEWS several communications on Flour Mill Explosions. In these reference is made to a letter of J. Lawrence Smith's to M. Dumas in respect to this matter. This letter I have not yet seen, but judging from the fact that Dr. Smith was supplied by myself with all of the information to be obtained here on the subject, including descriptions of experiments made before the coroner's jury by Professor L. W. Peck, a summary of the testimony given, and plans of the buildings destroyed, his opinions were based upon the facts.

In the *American Journal of Science and Arts* for October, 1878, will be found an article in which I made a record of all of the facts that I had learned likely to prove of scientific interest, to which I beg the attention of any one interested. Professor Peck and myself were requested to attend the sittings of the coroner's jury to act as experts and examine witnesses. We heard every word of the testimony, and very soon agreed in the conclusion that the mills exploded from ignition of a mixture of wheat dust and air. In fact, we came to that conclusion before we heard the testimony, and within twenty-four hours of the time the mills exploded. The testimony only established these convictions, and the experimental and theoretical proof placed them beyond doubt or question. In the controversy that followed the publication of these opinions, we for the first time saw a statement of the views held by Professor McAdam, and extracts from his paper. It appears to me that no other conclusion is possible when the subject is discussed on scientific principles, and that all of the talk about "distilling flour," "hydrogen," and "mystery," is superlative nonsense.

I enclose by the same mail a copy of the reports made by Professor Peck and myself before the coroner's jury, also the paper above referred to from the *American Journal of Science and Arts* and a paper from the *Popular Science Monthly*, in which the experiments are described by which the explosive force generated by burning flour dust is shown. It is, of course, too much to ask you to re-publish these; but on looking them over you may conclude to make such extracts as will assist in disseminating correct information on a subject of so much public importance. Both English and American millers are slow to believe that a flour sack can contain so much potential energy, yet we have here among those who produce the weekly grist of 35,000 lbs., men of long experience, who wonder that any intelligent person should question such a statement.—I am, &c.,

S. F. PECKHAM.

July 16, 1879.

PECULIAR SPECTRA OF LIGHTNING.

To the Editor of the Chemical News.

SIR.—In your issue of August, 25th, 1871 (CHEMICAL NEWS, vol. xxiv, p. 96), I endeavoured to describe some peculiar spectra of lightning observed in Cornwall, and which were kaleidoscopic in their complexity; the dominant spectra, however, being suggestive of copper. Subsequent enquiry and reflection made it probable that the spectra were really those of copper and of other metals; for in the general direction in which I examined the lightning was a smelting-house, from the tall chimney of

which issued the fumes arising from the processes connected with the extraction of silver from lead and copper ores; and on the evening of my observation a remarkably brilliant display of varied colour had been observed accompanying the play of the lightning upon the metallic and metalliferous heaps lying around the yard of the works. Nor should it be forgotten that the district itself is a metalliferous one.

Numerous examinations since then of the spectra of lightning in two other counties—Sussex and Hants—have revealed to me no similar spectra in the atmosphere.

The intense fusing power of atmospheric electric discharges upon the mineral substances of the earth, as instanced in the "fulgurites" or silicious tubes penetrating the sands of South America and other places, suggested to me that another but transient effect of the discharge would be the diffusion along its path of metallic substances, strikingly contrasted in scale with metallic diffusion along the voltaic arc of the laboratory. Gold, from its occurrence in the metallic state, ought thus readily to reveal its situation, whether in regions easily accessible or not. It seems obvious that the instrumental observation of the spectroscopic character, and the direction of the lightning with the interval of flash and report, would be sufficient to determine the character and situation of exposed metalliferous deposits in unexplored districts. Nowhere would such observations be better repaid, perhaps, than in Natal; for as recorded by Dr. Mann, nowhere in the world can the colorific displays of lightning be grander than in Natal. Indeed, my somewhat old ideas on the subject have been revived by our present interest in that quarter.—I am, &c.,

JOSEPH GIBBONS.

Uphurstbourne, Hants, July 31, 1879.

DETERMINATION OF SILICON IN IRON AND STEEL.

To the Editor of the Chemical News.

SIR.—The experiments of Mr. Thomas M. Brown, described in the CHEMICAL NEWS for July 25th (page 40), are interesting and of practical value; but there is one assumption which is not warranted by the facts.

Mr. Brown, and many other chemists, assume that when silicious iron is dissolved in hydrochloric or sulphuric acid, the white residue which separates consists of silica, and Mr. Brown remarks that the silica resulting from the action of either of the above acids is "light and flaky," while nitric acid gives a compact and granular product.

It has been shown by several chemists, and attention was drawn to the fact by me in a letter which appeared in the CHEMICAL NEWS for February 20, 1874, that the light flocculent residue obtained by the solution of silicious iron in hydrochloric or sulphuric acid consists chiefly of a lower oxide of silicon (called by Wöhler "Leukon"), the formula of which has been differently stated as $2\text{SiO}, \text{H}_2\text{O}$, and $\text{SiO}, \text{H}_2\text{O}$. The latter formula corresponds to that of a silico-formic acid, $\text{H}, \text{SiO}, \text{OH}$. At any rate, the fact that the body in question is a less highly oxidised compound of silicon than SiO_2 is proved by the following reactions.—

1. The residue dissolves in alkalis with evolution of hydrogen and formation of an alkaline silicate.
2. The washed residue is not wholly insoluble in water, and the aqueous solution is a powerful reducing agent.

Evidently, then, the proportion of silicon which passes into solution is dependent on the volume of liquid present and the quantity of water used for washing. No doubt the concentration of the acid and the temperature of the operation are also important factors, and, in short, the number of the conditions affecting the result is so large that the proportion of silicon which passes into solution is little other than accidental. As the filtrate contains the lower oxide and not silica, it does not at all follow that simple evaporation to dryness is sufficient to render it

insoluble. However, during evaporation more or less oxidation occurs, and if the solution contain any ferric chloride there can be little doubt that the whole of the dissolved silicon is recovered as insoluble silica.

I may add that I have several times had reason to suspect the formation of traces of silicide of hydrogen during the solution of highly silicious spiegeleisen in hydrochloric acid, but have never succeeded in verifying my suspicions.

I note that Mr. Brown finds that the "silica" obtained by the solution of the iron in sulphuric acid will seldom burn white. This is in accordance with the known properties of the lower oxide of silicon, which on ignition gives off silicide of hydrogen, and this is decomposed with deposition of more or less silicon.

In conclusion, I may say that five years' additional experience has resulted in still stronger confidence in the method of estimating silicon described by me in the *CHEMICAL NEWS*, vol. xxix, p. 91.—I am, &c.,

ALFRED H. ALLEN.

Sheffield, August 6th, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 26, June 30, 1879.

Chemical Constitution of Alkaline Amalgams.—M. Berthelot.—The author has prepared a series of alkaline amalgams, both liquid and solid, treated them with dilute hydrochloric acid and measured the heat evolved. At the same time the analysis of the resulting liquid showed the proportion of alkaline metal contained in each, which is always less than that employed in the preparation. The heat of oxidation of amalgams rich in potassium exceeds that of amalgams rich in sodium. It is not the same with the amalgams richest in mercury, the heat of formation of such potassium amalgams surpassing that of the corresponding sodium amalgams by a quantity which may reach +8.6 cal. for $Hg_{12}K$ as compared with $Hg_{12}Na$. The oxidation heat of potassium surpasses that of sodium in an inverse direction and only by +4.7 cal. Hence the oxidation heat of potassium amalgam may be reduced to +4.8 cal., that of sodium in analogous conditions being +5.6 cal. This accounts for the displacement of potassium in a solution of potassa by sodium amalgam.

Peculiarity of an Experiment by Gay-Lussac and Thénard.—H. Debray.—These chemists in their experiments on the preparation of alkaline metals passed the hydrates of potassa or of soda in vapour over an excess of iron contained in a gun-barrel heated to the highest possible temperature. Hydrogen and vapours of potassium and sodium escape from the apparatus, and the corresponding oxygen remains fixed upon a part of the iron, but chiefly the part outside the furnace and relatively the least hot. The iron is in great excess in proportion to the quantity of oxygen to be fixed; hence, whatever may be the internal reactions produced at elevated temperatures it is fixed upon the interior parts.

Spectral Examination of Ytterbia.—M. Lecoq de Boisbaudran.—The author has submitted the aqueous solution of the chloride to the induction spark, and has thus obtained a fine spectrum, formed chiefly of bands grouped between the solar rays D and E. These bands are almost all shaded from the left to the right. The author thinks the existence of a specific emission spectrum for this new element the more important as its chemical reactions are not well characterised.

The Law of Stokes, in Reply to M. E. Becquerel.—S. Lamansky.—The author describes experiments to show that the difference between the phenomena of phosphorescence and of fluorescence is not confined to a difference in the duration of the luminous emission after the previous influence of the exciting rays.

Dissociation of Ammonium Hydrosulphate.—R. Engel and A. Moitessier.—A reply to the paper of M. H. Sainte-Claire Deville, of June 9.

Action of Phthalic Anhydride upon Naphthalin in Presence of Ammonium Chloride.—E. Ador and J. M. Crafts.—By this reaction the authors have obtained naphthoyl-orthobenzoic acid.

Biedermann's Central-blatt.

July, 1879.

Ammoniacal Salts in the Seas of the Present and of Former Times.—L. Dieulafoy.—The presence of ammonia in sea-water is general. It varies in quantity from 0.2 to 0.36 m.grm. The deposits from concentrated sea-water are strongly ammoniacal, that of the Pool of Lavalduc containing 250 times as much ammonia as the water of the Seine during its passage through Paris. Ammonia is present to a notable extent in gypsum, which must be regarded as a sedimentary formation from former seas.

Distribution of Ammonia.—Dr. Angus Smith.—From the *CHEMICAL NEWS*.

Contributions to the Establishment of a Rational System of Examining Water as regards its Influence upon the Health of Men and Animals.—Dr. F. Holdeiss.—See page 63.

Influence of a Stratum of Vegetation and of Shade upon the Physical Properties and the Fertility of the Soil.—Prof. E. Woolny.—This paper does not admit of useful abstraction.

Absorptive Power of the Constituents of the Soil for Gases.—G. Ammon.—The condensing power of the ingredients examined may be arranged thus for watery vapour and ammonia:—Ferric oxide, humus, gypsum, kaolin, carbonate of lime. The action is greatest between the temperatures 0° and +10°, and decreases progressively both above and below these points. Hydrated ferric oxide converts the larger portion of the absorbed ammonia into nitric acid.

Justus Liebig's Annalen der Chemie,
Band 197, Heft 3.

Amidines and Thiamides of Monobasic Organic Acids.—A. Bernthsen.—The author describes here the ethers of the thiamidic acids.

The Composition of Cinchonin.—Zd. H. Skraup.—The author's analytical results agree with the formula $C_{19}H_{21}N_3O$, better than with that generally accepted, $C_{20}H_{24}N_3O$. He also examines a number of its salts.

Oxidation Products of Cinchonin.—Zd. H. Skraup.—The author examines the production of cinchotenin by the action of potassic permanganate upon cinchonin. A volatile acid was obtained which proved to be pure formic acid.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 5, 1879.

On Antichlor.—G. Lunge.—The author shows that a molecule of hyposulphite does not suffice for an atom of chlorine. Ammonia, which was proposed by Kolb as an antichlor, acts very imperfectly in dilute solutions and at ordinary temperatures, and the pungent odour produced by the reaction of ammonia or ammoniacal salts with the hypochlorites renders its practical application as an antichlor inadmissible.

Organic Ferrocyanides.—O. Bernheimer.—An account of the ferricyanides of tetra-methyl-ammonium and tetra-ethyl-ammonium.

On Berberin.—H. Weidel.—On treating berberin with nitric acid the author obtains no oxalic acid, but a finely-crystalline body, which he names berberonic acid, the preparation, properties, &c., of which are described at length.

Action of Melting Caustic Soda upon Phenol and Synthesis of Phloro-glucin.—L. Barth and J. Schreder.—The principal products of the reaction are pyrocatechin, resorcin, and phloroglucin.

On Cinchonin.—M. Fileti.—A preliminary communication. The author has obtained novel compounds on saturating a dilute hydrochloric solution of cinchonin with chlorine and bromine respectively.

Conquinin Sulphate.—O. Hesse.—A solution of conquinin sulphate in chloroform after standing for some months displays a green fluorescence almost as strong as Schær's solution of quinidin sulphate in chloroform.

Remarks on the Treatise of MM. Weselsky and Benedikt, "On certain Azo Compounds" (xii., 226).—P. Griess.—The author claims priority as regards the discovery of azo-benzol-naphthyl-amin. He considers that the practical application of such compounds as dyes is hindered by their sparing solubility in water, though their sulphacids may be applicable as orange dyes. If a liquid containing traces of nitrous acid is acidulated with pure sulphuric acid, then mixed with some solution of sulphanilic acid, and in about ten minutes afterwards with a few drops of a solution of naphthylamin sulphate, previously decolourised with animal charcoal, a red colouration shortly appears. In this manner nitrous acid can be detected in human saliva. In urine, contrary to the assertion of Schönbein, it does not appear to be present.

On Isocymol.—O. Jacobsen.—The author describes a number of derivatives of this compound.

Oxyparaxylylic Acid.—O. Jacobsen.—An account of certain salts of this acid.

Purification of Mercury.—L. Meyer.—The author applies a dilute solution of ferric chloride in an apparatus which he here figures.

The Relation between the Melting-points of the Elements and their Coefficients of Expansion by Heat.—T. Carnelley.—A table which the author has compiled shows that with five exceptions the coefficient of expansion increases as the melting-point sinks. The five exceptions are arsenic, antimony, bismuth, tellurium, and tin; the three former of which belong to the same elementary group, and even these among themselves display a similar relation between their melting-points and coefficients of expansion. Why these five bodies form an exception does not yet appear. It must, however, be noted that they are all found on the ascending side of Meyer's Curve of the Elements (see his "Modern Theories of Chemistry"), whilst three of them, tin, antimony, and tellurium, follow immediately upon each other in the above curve. Arsenic, antimony, bismuth, and tellurium are all very brittle and belong to the same crystalline system, and bismuth and antimony are the only two known pure elementary bodies which expand on congealing. Tin, in some of its compounds, displays an abnormal melting-point as will be shown in a future memoir. Both the melting-points and the coefficients of expansion may be periodic functions of the atomic weight.

Presence of Meta-nitro-toluol in Commercial Nitro-toluol.—P. Monnet, F. Reverdin, and E. Nölting.—The authors have detected meta-toluydin by examining the acetyl derivatives of a commercial toluydin, free from aniline.

Part Played by Meta-toluydin in the Formation of Magenta.—P. Monnet, F. Reverdin, and E. Nölting.—The presence of meta-toluydin is in every mixture injurious to the purity of tone of the dye produced.

Quantitative Determination of Zinc.—F. Beilstein and L. Jawein.—The solution of the zinc, nitric or sulphuric, is mixed with soda till a precipitate is formed, and then with potassium cyanide till a clear solution remains. Platinum electrodes, conveying the current of four Bunsen's elements, are plunged into the liquid, which is kept cool if needful by plunging the beaker in a vessel of water. When all the zinc appears to be precipitated the electrodes are lifted out of the liquid, and the zinc is washed, first with water, then with alcohol, lastly with ether, and dried in an exsiccator. After weighing it is dissolved in hydrochloric or nitric acid, and the electrode, cleansed and weighed, is re-introduced into the liquid to ascertain whether the zinc has been completely precipitated.

Treatment of Bunsen Elements.—F. Beilstein and L. Jawein.—To preserve connecting wires, binding screws, &c., from rusting, the authors recommend that when cleaned they should be anointed with oleo-naphtha, a lubricant obtained from Caucasian petroleum.

On Nitrochinolin.—W. Königs.—An examination of nitrochinolin, amidochinolin, and amidolepidin.

Modification of Simpson's Method of Determining Nitrogen.—W. Hankó.—This paper requires the accompanying engraving.

Synthesis of Chinolin from Allyl-anilin.—W. Königs.—Allyl-anilin is passed over lead oxide at low redness, when chinolin is found among the products.

Production of Nickel and Cobalt in a State capable of being Rolled.—T. Fleitmann.—The author conceived the idea that the brittleness of these metals was due to absorption of carbonic oxide, and succeeded in removing it by the addition of small quantities of magnesium, obtaining both metals in a perfectly malleable and ductile condition. The magnesium is introduced through a hole in the lid of the crucible after the oxygen has been removed by the addition of fragments of charcoal. Otherwise explosions may ensue.

Action of Phosphorus Penta-chloride upon Isatin and Allied Bodies.—A. Baeyer.—The author considers that the best method of obtaining indigotin from isatin chloride is to treat the latter with a solution of hydriodic acid in glacial acetic acid.

Amido-nitro-sulphide of Iron.—W. Demel.—This somewhat complex body is obtained by the reaction of ammonium sulphide and potassium nitrite with the solution of a ferrous salt.

Novel Method of Forming Ketons.—G. v. Bechi.—Freund discovered a general method for the formation of ketons by the action of acid chlorides upon zinc-alkyles. The author finds that iodalkyles in presence of metallic sodium occasion the same reaction.

Proportion of Water in Calcium Glycolate.—C. Böttiger.—This salt crystallises with variable proportions of water according to temperature, &c.

Contribution to a Knowledge of the Ureids: Synthesis of Dimethyl-barbituric Acid.—E. Mulder.—Not susceptible of useful abstraction.

Oxidation of Ortho-toluol-sulphamids.—O. Fahlberg and Ira Remsen.—Not suitable for abstraction.

Schizomycetous Fermentation.—A. Fitz.—The principal fact here established is the non-existence of the so-called butyro-acetic acid.

Azo, Azoxy, and Hydrazo Compounds.—H. Schmidt and G. Schultz.—With the removal of oxygen from the nitro compounds the melting-point rises up to the azo compounds; the hydrazo compounds melt at a lower temperature than the azo and even the azoxy compounds (except hydrazo-benzol); and the amido derivatives melt lower than the corresponding nitro compounds.

On Diphenyl Bases.—H. Schmidt and G. Schultz.

On Diphenoles.—H. Schmidt and G. Schultz.—These two memoirs do not admit of useful abstraction.

No. 6, 1879.

Oxidation of Resorcin to Phloroglucin.—L. Barth and J. Schreder.—The operation is effected by melting resorcin in a silver capsule with a considerable excess of caustic soda. Shortly after the development of gases has decreased the source of heat is removed, the mass when cold is dissolved in dilute sulphuric acid, the liquid filtered, and the filtrate repeatedly extracted with ether. Phloroglucin crystallises from the ethereal solution.

Formation of Ozone by Hydrocarbons.—J. Schiel.—It has been observed that the alkaline metals, thallium, &c., do not retain their brightness if kept in a stoppered bottle under rock oil. The author proves that this action is due to the formation of ozone.

On Fermentation.—J. Schiel.—The author passed the current of two zinc carbon elements through a solution of sugar, mixed with yeast, juice of meat, and a little ammonium phosphate. The fermentation proceeded as usual, but the formation of bacteria was prevented.

Action of Nitrous Acid upon Tin-phenyl-chloride.—B. Aronheim.—The compound obtained was identical with tin-triphenyl-chloride. Nitroso-benzol was also produced.

Action of Gaseous Chlorine upon the Hydrates of Baryta and Strontia (II.).—J. Konigels-Weisberg.—In this memoir the author describes the action of chlorine upon hydrate of strontia containing different proportions of water.

Action of Nitrous Anhydride upon Proto-catechuic Acid.—Max Gruber.—The products obtained were oxalic and picric acids, carboxy-tartronic acid, dinitro-dioxy-quinon, α -dinitro-phenol, and nitro oxy-benzoic acid.

Methyl Derivatives of Para-phenylen-diamin.—C. Wurster.—An examination of dimethyl-para-phenylen-diamin, its acetyl derivative, and of the tetra-methyl compound.

On Nitro-dimethyl-anilin.—C. Wurster.—Nitroso-dimethyl-anilin is readily converted into nitro-dimethyl-anilin by the action of oxidising agents. The author confirms Weber's account of its preparation.

Ureas of Dimethyl-para-phenylen-diamin.—F. Binder.—The author has obtained and described two ureas, a mono and a di compound.

MISCELLANEOUS.

University of London.—The following are lists of the candidates who have passed the recent B.Sc. examination:—First B.Sc. Examination: Pass List (*First Division*).—T. J. Bowlker, private study; J. S. W. Chitty, Magdalen College, Oxford; T. R. Dallmeyer, University College and private study; G. L. V. Fairbault, Catholic University College, Kensington; P. F. Frankland, Royal School of Mines and private study; E. T. Glasspool, M.A., private study; W. L. Goodwin, University of Edinburgh; C. Grimshaw, Owens College and private tuition; S. F. Harmer, University College; G. W. Hill, King's College and St. George's Hospital; W. Law, private study; R. B. Lee, private study; C. Myhill, University College and private tuition; W. Odling, Royal School of Mines; F. W. Payne, B.A., Brighton Grammar School; W. J. C. Ross, King's College and private study; J. Ryan, Unattached, Cambridge; G. P. Simpson, private tuition; A. Smithells, Owens College; A. K. A. Spiegel, Owens College; A. F. Stoddart, University College, Bristol; Ellen Martha Watson, private study; S. Young, Owens College. (*Second Division*).—W. Bryant, B.A., private study; R. Frost, Owens College; J. R. Green, private study; Catherine Alice Raisin, private study; G. A. T. Walton, St. Bartholomew's Hospital; H. Wilson, The Leys School, Cambridge. (*Mathematics only*).—W. J.

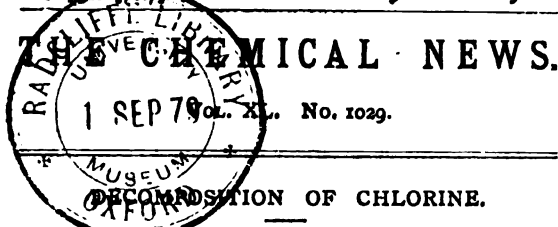
Collins, St. Bartholomew's Hospital; A. Newsholme, St. Thomas's Hospital.

Letters Patent for Colouring Matters.—A patent case of some interest was argued before the Lord Chancellor on the 30th ult. Mr. Frank Wirth, of Frankfort, petitioned that the seal might be affixed to letters patent for an invention for "Improvements in colouring matters, and in the manufacture of the same," communicated to him by Messrs. Meister, Lucius, and Brüning, of Höchst, Germany. It appeared that the improvements consisted substantially in the preparation of colouring matters by the reaction of diazo compounds of phenols upon the disulpho acids of beta-naphthol. The application was opposed by Dr. J. P. Griess, F.R.S., of Burton-on-Trent, chiefly on the ground that the applicant's invention was already covered by previous letters patent granted to him, the opponent. These letters patent were respectively dated the 4th of October, 1877, No. 3698, and the 20th of November, 1878, No. 4726, and it appeared, from the specifications of these letters patent, that Dr. Griess had, in fact, patented the preparation of colouring matters by acting upon the diazo compounds of phenols and phenolic ethers with, among other things, disulpho acid of beta-naphthol. It was, however, pointed out on behalf of the applicant that the latter did not claim to use the disulpho acid of beta-naphthol as used by Dr. Griess under his letters patent, but one or other of the two isomeric disulpho acids of beta-naphthol which Messrs. Meister, Lucius, and Brüning had been the first to discover, and the separation of which and their respective application or treatment with certain other substances, such as toluydin and xylochin, Mr. Wirth had been the first to patent, that is to say, on the 29th April, 1878, No. 1715. It appeared that these isomeric disulpho acids were capable of reactions different from those resulting from the use of the disulpho acid described in the specifications of Dr. Griess. The application for letters patent by Mr. Wirth had been accompanied by the usual statutory declaration, which, however, had been declared at Frankfort before H.B.M. Consul. The objections filed on behalf of Dr. Griess were in substance (1) that F. Wirth was not the true and first inventor or importer of the alleged invention, and (2) that the alleged invention was wholly or in part covered by the letters patent of Griess. By his affidavit in opposition the latter had also objected that Wirth had not made and could not make at Frankfort a declaration that would be good under British patent or other law. There had been no opposition before the law officer. Upon the petition being called on, Mr. Aston, Q.C., urged as a preliminary objection that Wirth's application was void *ab initio*, on the ground that a declaration such as that required by the Patent Law Amendment Act could not be made by a foreigner resident abroad before the British Consul, contending that the statutory forms were not merely directory but imperative. Moreover, Wirth, being a resident abroad, could not receive a communication from other foreigners also abroad, and receive a British patent as for an imported invention. The Lord Chancellor was of opinion that letters patent might be granted to a foreigner resident abroad for an invention communicated to him by another foreigner also resident abroad. The petition was then opened and heard upon the merits. Affidavits by Professor Odling, F.R.S., and Professor Armstrong, F.R.S., were read on behalf of the opponent, and ultimately it was arranged that the letters patent of the applicant should be sealed upon his undertaking to insert in his specification a disclaimer of any matters covered by the letters patent granted to Dr. Griess, and no order was made as to costs.

TO CORRESPONDENTS.

W. J. Reynolds.—Mix the paint with gelatine and water instead of oil.

M. T. D.—The address was printed just as it was received from Messrs. W. H. Smith and Son.



SINCE the publication of Prof. Meyer's important researches on chlorine in the CHEMICAL NEWS of August 1, we have obtained some additional particulars, which we hasten to lay before our readers. The vapour-density apparatus designed by Prof. Meyer, and referred to by Mr. Watson Smith, has already been shown in the CHEMICAL NEWS (vol. xxxix., p. 66), and it was by means of this apparatus, suspended in a Perrot's furnace, we believe, that the oxygen was obtained. In his note Mr. Watson Smith said that, "Experimental evidence appears to indicate beyond a doubt that to all appearance (*"Moglicherweise"* is the word Victor Meyer wishes at present to be used) by heating chlorine oxygen can be obtained." The chlorine was prepared absolutely pure by heating PtCl_4 (or Pt_2Cl_6), and was dried with every precaution, being passed over CaCl_2 , through SO_4H_2 , and over a layer of P_2O_5 one metre in length; it was then passed into the vapour-density apparatus, in the lower portion of which porcelain was substituted for glass. The dissociated chlorine gas issuing out of the small tube *a* and containing oxygen was absorbed by passing through KI solution. The oxygen passing through unabsorbed was collected, and the usual tests were applied.

SYNTHESIS OF PHENYL-NAPHTHALENE.

By WATSON SMITH, F.C.S., F.L.C.

In my last papers, or rather preliminary notes, on the above subject (see CHEMICAL NEWS, vol. xl., p. 3; also *Ber. der Deutsch. Chem. Gesell.*, xii., p. 1396) I described a method by which iso-dinaphthyl, diphenyl, and a new hydrocarbon—probably phenyl-naphthalene—were formed together, and could readily be separated by fractional distillation, and by subsequent treatment of the crude new substance with boiling dilute spirits of wine the new hydrocarbon was dissolved out, but the iso-dinaphthyl mostly remained insoluble. On sublimation the new body was easily obtained quite pure, and formed beautiful small plates with blue fluorescence, the vapours possessing a most pleasant odour of oranges. In the above case the calculated quantities of brom-benzene and naphthalene were passed through the red-hot tube. The experiment was now conducted differently, soda-lime being introduced into the tube instead of pumice-stone. It was now found that the reaction proceeded much less favourably, and that benzene was formed and came over, no doubt with a certain quantity of diphenyl. This reaction is analogous to that previously described (CHEMICAL NEWS, vol. xl., p. 3), in which brom-naphthalene and benzene were passed over soda-lime, and naphthalene was formed. The former method was therefore adhered to, but I determined to modify the proportions, so as, if possible, to make use of the tendency to form diphenyl so as to yield me rather an increased yield of phenyl-naphthalene. This was effected by employing in the next mixture used a certain excess of naphthalene over the calculated quantity. The soft tarry-looking mass was then distilled, and the unaltered brom-benzene and naphthalene coming over, after being mixed with more naphthalene, were passed again through the red-hot tube. This was continued till but little undecomposed brom-benzene passed over, or till it was almost all decomposed. On subjecting the mixture of high

boiling residues thus obtained to distillation, unaltered naphthalin, very little diphenyl, a greatly increased quantity of phenyl-naphthalene, and iso-dinaphthyl, with still higher boiling products, came over. The amount of phenyl-naphthalene now obtained was equal to, if not larger than, that of the dinaphthyl formed—a vast improvement on the results of the former methods.

A combustion was now made of the purified substance, with the following results:—

	Found.	Calculated for C_{10}H_7 C_6H_5
Carbon	94.63	94.12
Hydrogen	6.01	5.88

Two vapour-density determinations were also made according to the method of Victor Meyer and Carl Meyer, and these were conducted conjointly with my friend Carl Meyer with the utmost care and exactitude. We made the determinations in an atmosphere of nitrogen, and obtained the following results:—

I.	II.
$B = 722.8$ $w = 24.3$ $t = 25.5$ $S = 0.1336$ $V = 17.3 \text{ c.c.}$	$B = 722.8$ $w = 24.3$ $t = 25.5$ $S = 0.13825$ $V = 17.8 \text{ c.c.}$
$d = \frac{S(1 + 0.003665 \cdot t) 587780}{(B - w)V}$	

Hence—

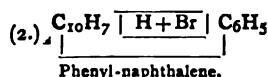
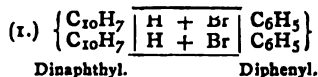
	Vapour-density.
	Found. Cal. for C_{10}H_7 C_6H_5
(I.) . . .	$\frac{0.1336(1 + 0.003665 \cdot 25.5) 587780}{(722.8 - 24.3) 17.3} = 7.10$
(II.) . .	$\frac{0.13825(1 + 0.003665 \cdot 25.5) 587780}{722.8 - 24.3) 17.3} = 7.14$

The melting-point was now re-determined with great care, using one of Giessler's normal thermometers; it was found to be 95° as the mean of several determinations, and this must be regarded, then, as the corrected melting-point, that first given (101° to 102°) having been determined with an ordinary instrument. This latter thermometer, with the freshly-prepared substance, indicated 100° to 101° ; hence showing that a substance practically of a constant composition—in fact, the same body—had been obtained as that tested in the first case.

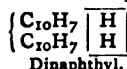
A combustion was now made again, but of the very substances which had been the subjects of the two vapour-density determinations. The contents of the glass vapour-density apparatus (Meyer's), were treated with hot absolute alcohol, well shaken, and then thrown on to a filter. The fine white sand (used to break the fall of the little tube with substance), remained, and the alcoholic solution of the phenyl-naphthalene passing through, was diluted with much water, which precipitated the phenyl-naphthalene in small microscopic and pure white plates. This, after washing with water, was dried at 70° in the air-bath, and analysed. 0.23925 grm. substance gave 0.1185 water and 0.82795 grm. carbon dioxide:—

	Found.	Calculated.
Carbon	94.37	94.12
Hydrogen	5.51	5.88
	99.88	100.00

In the process of preparation it would seem, then, that when the calculated quantities of brom-benzene and naphthalene are taken, mixed, and passed once through the red-hot tube, the following reactions take place together:—

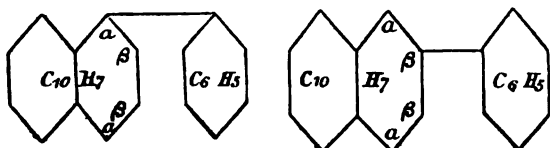


But on using an excess of naphthalene, and passing the undecomposed brom-benzene and naphthalene, with the addition of a little more naphthalene, through the red-hot tube till all brom-benzene disappears, it would seem that reaction (1) ceases to a considerable extent, and for the most part, in fact, reaction (2) now predominating, together with the simpler one,—

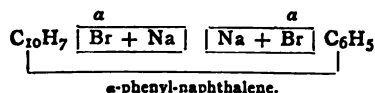


resulting by the action of heat on the excess of naphthalene, and hence accounting for the presence in the reaction-product of so considerable a quantity of iso-dinaphthyl, with scarcely any diphenyl.

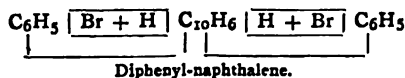
There are two isomerides of phenyl-naphthalene possible, viz., the α and the β :—



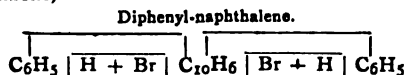
It is interesting to ascertain which isomer represents the new hydrocarbon obtained. I think there can be little doubt it is the β -phenyl-naphthalene, and for the following reason. It is formed from that naphthyl residue of a naphthalene group, which is itself formed at a high temperature the most readily and easily. Now, by the action of heat alone, or heat in conjunction with halogenic action, if the term may be allowed, upon naphthalene, my previous investigations have shown that iso-dinaphthyl is formed in preponderating quantity, and that iso-dinaphthyl is in the highest degree of probability the β β -dinaphthyl, hence the phenyl-naphthalene is in all probability the β -isomeride. It is not at all improbable the α -isomeride may be obtainable by means of Fittig's reaction, with a mixture of α -brom-naphthalene and α -brom-benzene; thus—



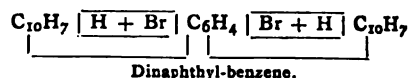
This question I shall next proceed to solve, together with the following :—"Is it not possible to obtain a diphenyl-naphthalene by the action, say, of two molecules of brom-benzene on one of naphthalene on passing the mixture through the red-hot tube?" Thus—



or by the action of 2 mols. benzene on 1 of dibrom-naphthalene,—



Or it is possible, by using a mixture of 1 mol. dibrom-benzene and 2 mols. of naphthalene, that dinaphthyl-benzene may be obtained :—



Indeed, I am proceeding to examine the higher boiling-products obtained, with the dinaphthyl and phenyl-naphthalene, when excess of naphthalene was used, to see if small quantities, at least, of the above hydrocarbon be not already present. The solution of these latter problems will form a considerable portion of my future work.

University Laboratory, Zürich,
August 4, 1879.

UPON AMMONIUM NITRITE, AND UPON THE BY-PRODUCTS OBTAINED IN THE OZONISATION OF AIR BY MOIST PHOSPHORUS.

By ALBERT R. LEEDS.

As long ago as 1848, Dr. T. Sterry Hunt threw out the suggestion, that the nitrogen of the atmosphere is really composed of two equivalents of the element, sustaining towards each other the same relations as the two equivalents in nitrous oxide. He supposed that the group NNO is not a simple oxide, but an anhydrous amide, or nitryl, derived from the ammonium nitrate by the removal of 2H₂O, and capable, when passed over a mixture of lime and potash at a sufficiently high temperature, of regenerating ammonia and a nitrate.* He insisted upon the parallelism between this case and that of ammonium nitrite which undergoes a precisely similar decomposition under the influence of heat, losing 2H₂O and evolving NN, nitrogen gas. It would appear that Prof. G. C. Schaeffer had independently arrived at a similar conclusion concerning the dual nature of nitrogen, holding the same view as Dr. Hunt, that it is the nitryl of ammonium nitrite, and capable of forming this body by assuming again the elements of water.†

In 1862, Schönbein published an extensive series of experiments upon the generation of ammonium nitrite from water and atmospheric air under the influence of heat; Unacquainted, apparently, with speculations by which the formation of ammonium nitrite in this manner had been anticipated on theoretical grounds, he announced his own discovery as in the highest degree remarkable and hitherto entirely unexpected. He states that his own labours in this direction were animated by the consideration that this salt is so readily decomposed under the influence of heat, into water and nitrogen, that it might, under suitable circumstances, be likewise readily regenerated from the same bodies. These circumstances he found to consist in the contact of water, converted into vapour at various temperatures, with atmospheric air, and attributed the combination principally to the heat employed. No other phenomena, according to Schönbein, were concerned in the generation of ammonium nitrite under these circumstances, and those which regulate its formation when various bodies, like hydrogen, hydrocarbons, fats, wood, coal, &c., were burned in air.

It is not necessary to enter into the details of his experiments, further than to remark, that in none of them, apparently, was the precaution taken to use air which had been purified from its pre-existent ammoniacal and nitrous compounds. This manifest source of error was at once pointed out by Bohlig,‡ who never succeeded, in whatever way or at whatever temperatures, the experiment was conducted, in obtaining the reactions of ammonium nitrite when purified air was employed, but readily obtained

* *American Journ. Sci.*, May, 1848, p. 407.

† "Chemical and Geological Essays," T. S. Hunt, and Ed., p. 472.

‡ *Ann. der Chem. und Pharm.*, 1862, vol. cxxiv., p. 1.

§ *Ann. der Chem. und Pharm.*, 1863, vol. cxxv., p. 21.

them when, under like conditions, common air was used. In opposition to these results of Bohlrig, Prof. Liebig maintained in the same journal, the trustworthiness of Schönbein's observations, contending that the fact was sufficiently proven by the very numerous and varied experiments of Schönbein, and disregarding the valid objection, that in none of these same experiments had purified air been employed. Even to the present day the generation of ammonium nitrite from atmospheric air and water by the aid of heat, is taught in text-books as an admitted fact, and that, too, after Zabelin, in 1864,* and Carius,† ten years later, had repeated, with exceeding care, and verified Bohlrig's observations. In particular, Carius employed most elaborate precautions, using only air and water which had been most carefully purified. The water was evaporated both with a rapid and a slow change of atmosphere; at various temperatures, from the common to 100°; both alone and after addition of baryta, the baryta being devoid of nitrogen compounds; in contact with platinum spirals, and diffused over a great surface of purified cotton-wool—in no case was ammonium nitrite formed.‡

It is of the highest importance, therefore:

I. To exclude the possibility of the conversion of water under the influence of heat in contact with atmospheric air, from among the number of possible sources of ammonium nitrite and nitrous compounds.

II. In cases of rapid oxidation, like the combustion of H,§ hydrocarbons,|| fats, phosphorus,¶ and other bodies in the air, if it be true that ammonium nitrite is formed, irrespective of any nitrogen compounds pre-existent in the

atmosphere, the origin of this ammonium nitrite is to be looked for in other causes than the conjunction of atmospheric air and water-vapour under the influence of heat.*

III. The same remark applies if any ammonium nitrite is formed by the slow oxidation of phosphorus in contact with air and water. The very numerous and laborious experiments by which Schönbein appears to have established the fact of the formation of ammonium nitrite under these circumstances, have caused its universal acceptance.† Bohlrig, following Schönbein, attributed the production of ammonium nitrite during the slow oxidation of moist phosphorus, to the action of the ozone formed at the same time. Carius, by elaborate proofs, has demonstrated that ozone, which was prepared by electrolysis in such a manner as to be entirely free from nitrogen compounds, and even from hydrogen gas, was entirely devoid of action upon nitrogen in the presence of vapour of water. In order to make the experiments comparable to those in which ozone is liberated at the same time with a disengagement of heat, as in the case of the electrical discharge, the ozone was brought by Carius into contact with nitrogen and water-vapour at all temperatures short of those at which ozone undergoes dissociation into ordinary oxygen—in every case with negative results. The ozone prepared by electrolysis was preferred to that obtained by the electrical discharge, since, as both Soret‡ and Carius have shown, ordinary oxygen prepared from potassium chlorate, and containing from 1 to 2 per cent of nitrogen, yields, when ozonised by the electrical discharge, small but determinable amounts of nitrous acid. Still later, Berthelot§ has investigated the accuracy of the statement made by Schönbein,|| that ozone combines with free nitrogen, in presence of alkalies, at ordinary temperatures, to form nitrous compounds.

Schönbein found in 3000 litres of air, powerfully ozonised by phosphorus, then washed with water, the washings treated with lime water, and the resulting calcium compound decomposed by potassium carbonate, an amount of nitric acid corresponding to 5 grms. of nitre. Berthelot collected in a number of flasks over water, oxygen ozonised by the silent discharge, until the flasks were three-fourths filled. The surplus water was then allowed to run out completely, its place being supplied by atmospheric air, and some pure baryta water introduced in each flask. After standing over night, all traces of ozone had completely disappeared. But the baryta water contained no traces of nitrogen compounds; the wash-water did. Similar results were obtained with the air ozonised by phosphorus, and completely washed before contact with baryta water.

Quite independently of the work done by other observers, an extended series of experiments had been instituted upon the phenomena which accompanied the ozonisation of moist air by means of phosphorus. In the earlier trials, attention was limited to the question whether oxidised compounds of nitrogen were produced or no. Subsequently the research was made to include all other by-products. It was deemed important to purify and measure the air used and the ozone formed; to determine the amount of phosphorus consumed, and of phosphoric and phosphorous acid produced; and, in case they were really present and it were possible to estimate them, the amounts of nitrogen compounds, of hydrogen peroxide and ozone, remaining in solution in the jar- and wash-waters. Last it be thought needless to have re-investigated these points the following reasons for so doing may be mentioned:—

I. Preliminary trials had shown that, if formed at all,

* In the case of hydrogen, Zöllner and Grete have shown, by the burning of very large volumes of perfectly pure hydrogen in completely purified air, that small amounts of ammonium nitrite are produced, and have demonstrated the presence of ammonia and nitrous acids in the water formed in the course of the combustion, by many concordant and quite satisfactory tests.—(*Berichte der Deutsch. Chem. Gesell.*, 1877, vol. x., pp. 2, 144.)

† See Carius, *Ann. der Chem. und Pharm.*, 1874, vol. clxiv. p. 43.

‡ *Comptes Rendus*, lvi., 390.

§ *Comptes Rendus*, 1877, lxxiv., 61.

|| *Denkschrift über das Ozon* p. 16 Basel, October, 1849.

* *Ann. der Chem. und Pharm.*, cxxx., p. 82.

† *Ann. der Chem. und Pharm.*, clxiv., p. 1.

‡ NOTE.—Since the above was written a paper has been published by A. v. Losicke (*Arch. Pharm.* [3], 14, 54 to 58, and *Chemiker Zeitung*, No 9, 1879). Inasmuch as I have no access to either of these journals, I can only quote from the abstracts (*Journ. Chem. Soc.*, April, 1879, p. 298; *CHEMICAL NEWS*, vol. xxxix., p. 150). It is stated in the *Journ. Chem. Soc.* abstract, that "the author has corroborated Schönbein's statement, that the evaporation of water in air produces ammonium nitrite, and gives the results of experiments to determine the conditions of its formation. It is found that ammonium nitrite is always formed when water evaporates freely, and the lower the temperature, the larger is the quantity produced; but the formation is prevented if the evaporation takes place in a narrow-necked flask. In another series of experiments it was observed that 1 litre of water, evaporated to a small bulk, yielded ammonia equivalent to 0.148 part in 100,000 parts of water; 1 litre, evaporated to small bulk at 40° to 50°, yielded ammonia equivalent to 0.5823 part of nitric acid per 100,000 of water; and, lastly, 5 litres of water, allowed to evaporate spontaneously, yielded ammonia equivalent to 2.9608 nitric acid per 100,000 parts of water. This last experiment shows the influence that the evaporation of rain-water and dew can have on the nourishment of plants; and it has been found that if a leaf be moistened and allowed to dry, nitrous acid is produced, and that in dew from leaves, ammonium nitrite can be easily detected."

With reference to these observations it is to be remarked:—1st. In the absence of any reference to the essential precautions of excluding the access of nitrogenised atmospheric particles, atmospheric ammonia, and nitrous compounds, it is fair to infer that these precautions were not taken, and in that case, the experiments of Bohlrig, Zabelin, and Carius, which contradict the hypothesis of Schönbein, must be regarded as unshaken and conclusive. 2nd. The rise of temperature, especially under such circumstances as would establish a current of vapour from the surface of the liquid, would diminish the rate of absorption of nitrogenous bodies from the surrounding atmosphere. Cooling the evaporating liquids would operate in a two-fold manner; in the first place, by augmenting the coefficient of solubility of the atmospheric ammoniacal compounds; and, secondly, by greatly increasing the length of time required for the spontaneous evaporation of so large a bulk as 5 litres of water.

To get rid of all traces of ammonia upon the surfaces of vessels is a problem of almost insuperable difficulty so long as laboratory operations must be performed in common air. In this connection we would call to mind the experiments of Faraday (*Quart. Journ. Science*, xix., 16), who was greatly perplexed by obtaining ammonia, on heating hydrogen gas in contact with potassium and zinc. The vessels and substances had been purified, and every precaution exercised, which the almost matchless skill of that incomparable experimental philosopher could devise, and yet Faraday abandoned the research, confessing his inability to satisfactorily account for the appearance of ammonia under these circumstances. Additions to our knowledge since that time may render it not presumptuous to suggest the cause above given.

§ Saussure (*Annales de Chimie*, lxxi., 282).

|| Böttger (*Jahresb. der Phys. Vereins, zu Frankfurt a.M.*, 1860, 1862, p. 66).

† Schönbein (*Ann. der Chem. und Pharm.*, vol. cxvii., p. 6).

there were grounds for supposing that these products were comparable in minuteness to the nitrogen compounds pre-existent in the atmosphere.

II. Since the time of Schönbein, the experiments of Goppelsröder and Carius had shown that the ozonisation of ammonia would produce ammonium nitrite and nitrate and hydrogen peroxide.

III. We could not discover that in experiments previously made upon the ozonisation of air by phosphorus, the air had been previously purified.

IV. It was important to determine whether any ammonia or nitrous acid existed at the close of the operation, or whether they were completely oxidised to nitric acid. The explanations of the reactions would depend largely upon the determination of these factors.

The phosphorus ozonator (described in the *Journal of the American Chemical Society*, vol. i., p. 8) was employed throughout the whole course of this investigation. In the first series of experiments ordinary air drawn from outside the laboratory was used. The jars in some cases contained common distilled water, in other cases the bichromate mixture made as described in article alluded to. The ozonised air was aspirated through the kerite tubing, from the last bell, into a wash-bottle containing *aqua puriss.* (re-distilled and free from every trace of ammonia), and then into one or more Peligot tubes containing neutral solution of potassium iodide. Only on titration was the solution of potassium iodide made acid. The reason for this precaution is given in the *Journ. Amer. Chem. Soc.*, vol. i., p. 78, where it is shown that a stream of air or oxygen passing through an acidified solution of potassium iodide, greatly increases the amount of iodide set free. The nitric acid in the wash-bottle was determined by reduction with pig-iron, the precautions being employed which are stated (*Proceedings Amer. Chem. Soc.*, vol. ii., No. 4, 1878).

TABLE I.
Nitric Acid Formed with Non-purified Air.

Air Used. Litres.	Total Ozone in Mgram.	Total Ozone in C.c. of Air.	V. p.c. of Air.	Total HNO ₃ .	HNO ₃ P.c.
27 (over H ₂ O)	—	—	—	0'111	0'000318
108 "	—	—	—	0'444	0'000318
45 (over CrO ₃)	13'61	6'37	0'014	0'314	0'00054
85 (over H ₂ O)	77'07	36'09	0'039	0'333	0'000303
90 (over CrO ₃)	71'91	33'67	0'037	0'610	0'00054

In the following trials, the air was purified by aspiration through a tube one-half metre in length, packed with cotton-wool, then through three Peligot tubes, the first containing *aqua puriss.*, the second soda solution, the third sulphuric acid. The soda employed in this and other trials was some made from sodium, and which had been repeatedly proven to be free from nitrogenous bodies. The sulphuric acid yielded to meta-diamido-benzol no trace of nitrous acid, nor any ammonia on neutralisation and reduction. Ordinary distilled water was employed in the jars.

TABLE II.
Nitric Acid Formed with Purified Air (*Aqua Pur.*).

Air Used. Litres.	Total Ozone. Mgrams.	Total Ozone. C.c. of Air.	V. p.c. of Air.	Total HNO ₃ . Mgrams.	HNO ₃ P.c.
262 (over H ₂ O)	261'8	104'16	0'04	1'11	0'000328

In an experiment similarly arranged, except that hydrant water was used in the jars, 0'3964 grm. of ozone was produced. The wash-water contained 0'37 mgrm. of nitric acid, and 1'3 mgrms. of hydrogen peroxide. The latter would correspond to 0'00039 per cent of the total weight of the air used.

In the foregoing experiments, the percentage of nitric acid as compared with the weight of the air passed over, appears quite constant. This is true whether water or bichromate mixture was employed, although with the latter the percentage was somewhat higher. The ozone had been manufactured for a variety of purposes, and

these percentages were not calculated until after all the determinations were concluded, when an unexpected closeness of agreement between the results was developed. Although this agreement pointed to a constantly operating cause, yet, by reason of the extreme minuteness of the amounts of nitric acid, it was deemed safer provisionally to assume that the agreement was really due to a constant source of error. A scheme was therefore drawn up, which should include the sources of error and the substances to be determined to the best of our knowledge. As others may detect features in which our methods were at fault, or essential precautions were overlooked, this scheme is given below:—

I. Use of a measured amount of *aqua puriss.*, in the jars.

II. Complete straining and washing of the air by means of cotton-wool, *aqua puriss.*, NaOH and H₂SO₄, both shown to be free from nitrogen.

III. Weight of phosphorus cakes before and after the experiment.

IV. Amount of ammonia in the jar-water at close of experiment.

V. Also of the nitrites and nitrates, as determined by reduction.

VI. Amount of nitrous acid, as determined by meta-diamido-benzol.

VII. Amounts of phosphoric and phosphorous acids and of hydrogen peroxide in the jar-water.

VIII. Estimation of H₂N, (HNO₂ and HNO₃), HNO₂, H₃PO₄, H₂PO₃, and H₂O₂, in the wash-water after ozonising ("ozone wash-water").

IX. Similar estimations, except of the phosphorus compounds, in the solutions employed to wash the air ("air wash-water").

X. Measurement of total volumes of air used, and of ozone after its escape from the ozone wash-water.

In order to compensate for the increase of suction necessitated by so many wash-bottles, the standards and discs carrying the phosphorus were raised nearly to top of the bells.

This required the employment of 11 litres of *aqua puriss.* in the jars. After 420 litres of air had been aspirated the jar-water was poured into three tall cylinders, the precipitate, mostly lead phosphate, allowed to settle, and the water and precipitate analysed separately. The weight of the phosphorus cakes at beginning of the experiment was 111'646 grms., at the close 54'284 grms. The total amount of ozone which passed out of the ozone wash-bottle, was 0'8046 grm., or an average of 1'92 mgrms. ozone per litre of air, the temperature varying during the course of the six days consumed in the experiment between 18° and 21°. This would correspond to 71'29 grms. of phosphorus consumed for each gramme of ozone produced. Probably the consumption would be in a somewhat different ratio at the temperature of maximum evolution (24°), and with bichromate mixture in the jars.

Phosphoric and Phosphorous Acids.

The amount of phosphoric acid (H₃PO₄) existing in solution in the jar-water, was 178'02 grms.; in combination with the lead as triplumbic phosphate, 0'939 grm.; and the comparatively small amount carried over into the ozone wash-water was 0'252 grm., making a total of 179'24 grms.

The large amount of phosphoric acid is itself a valuable product, and capable either of being used directly or of being easily re-converted into phosphorus. Hence, its formation cannot be considered as a drawback upon this method of ozonising. The phosphorous acid in the jar-water was 2'57 grms., in the ozone wash-water, 0'013 grm.

Ammonia, Nitrous and Nitric Acids.

(a.) In Air Wash-water.—The entire ammonia derived from washing 420 litres of air, was 0'03 mgrm., corresponding to 6 parts in 100 million. No nitrous acid could be detected by means of meta-diamido-benzol; nitric acid not determined.

(b.) In Jar-water.—50 c.c. of the jar-water rendered alkaline with pure soda, and distilled in a purified retort, yielded 0.08 mgrm. H_3N , corresponding to 17.60 mgrms. ammonia in the 11 litres. A like portion, similarly treated, gave, on reduction with pig-iron, 0.14 mgrm. H_3N , or in the total jar-water, 0.0308 grm. Both NH_4NO_2 and NH_4NO_3 should yield on reduction twice the amount of ammonia which they give when the ammonia is determined directly. In the above determination, the amount which was found falls short of the theoretical by a difference quite within the errors of experiment.

Since phosphorous acid was contained in the jar-water, it became important to determine whether this body, when distilled in neutral or slightly alkaline solution, with pig-iron, would yield a distillate capable of effecting the Nessler reagent. Some phosphorous acid was therefore prepared from phosphorus trichloride, and the experiment tried, but the results obtained negatived the above hypothesis.

No nitrous acid was detected. The nitric acid, as deduced from the ammonia found directly and after reduction amounted to 0.0489 grm., or 0.0089 per cent of the total weight of the air used.

(c.) In Ozone Wash-water.—The ammonia obtained by direct distillation amounted to 0.075 mgrm.; by reduction, to 0.315 mgrm.; this would leave for the ammonia, corresponding to the nitric acid, 0.24 mgrm. (equivalent to 0.88 mgrm. HNO_3), a result three times greater than the ammonia corresponding to the base, instead of twice, as should have been.

No nitrous acid was found by the use of meta-diamidobenzol.

Hydrogen Peroxide.

The ozone wash-water contained 2.01 mgrm. hydrogen peroxide, or 0.00037 per cent of the total weight of the air passed over. The nitric acid contained in the ozone wash-water, was only 0.000162, or about one-half the percentage of the hydrogen peroxide.*

Final Experiment.—The failure to find in the jar-water an amount of ammonia, on reduction, which would exactly correspond to the nitric acid required to form with the basic ammonia, ammonium nitrate, cast some suspicion on the above results. A final experiment was therefore instituted, in order to settle positively whether any ammonium nitrite could be detected at the close of the operation, and whether all the nitric acid found existed in combination as ammonium nitrate.

Renewed precautions were therefore instituted to purify the air employed—two columns of glass beads saturated with H_2SO_4 being employed in addition to the wash-bottles mentioned in the former trials. The air was forced by a tromp through this long series of purifiers into the ozonator, and drawn out again through the wash-bottles used to wash and titrate the products, by means of a water air-pump. By this means, the phosphorus cakes were always kept immersed to the same depth in the jar-water. Sliding standards had been discarded in the improved form of ozonator now employed, and the cakes were supported on leaden discs which rested upon little leaden brackets, passing through holes drilled through the sides of the bells, at a small distance above their lower rims. 4850 c.c. *aqua puriss.* were used in the jars; the amount of air passed through the apparatus was 481 litres; the ozone discharged from the ozone wash-water, 0.924 grm. The following table presents a synopsis of the results obtained:—

* NOTE.—In Table I. the ammonia found in the ozone wash-waters by reduction is all calculated as HNO_3 . If we, in like manner, convert the total ammonia contained in the air (see Table III.) into HNO_3 , it will be found equal to 0.00088 per cent of the air drawn over; subtracting this from 0.0003 per cent, the amount given for P over H_2O in Table I., leaves 0.00058 per cent, and this, if regarded not as it was when the experiments were made, as HNO_3 , but as NH_4NO_2 , would give for the true percentage of HNO_3 in Table I., 0.000167 per cent.

TABLE III.

Ammonia:		Air Wash-water.			
		HNO_3 .	HNO_2 .	NH_4NO_2 .	NH_4NO_3 .
Before Reduction.	After.	Mgrm.	Mgrm.	Mgrm.	Mgrm.
0.10	0.15	0.065	0.104	0.092	0.132
Corresponding in 100 Million Parts.					
Parts.		Parts.	Parts.	Parts.	Parts.
16	None.	10	17	15	20
Jar-water.					
Mgrms.	Mgrms.	Mgrms.		Mgrms.	
(1.) 7.76					
(2.) 4.82	11.64	None.	21.57	None.	27.39
Ozone Wash-water.					
0.066	0.198	None.	0.49	None.	0.31

It will be noted that the ammonia washed out of the atmospheric air is somewhat in excess of that required to neutralise the nitric and nitrous acids, and it may be queried whether this excess existed in some other form of combination, as the carbonate.

The determination marked (1) of ammonia in the jar-water, was made upon 50 c.c. Subsequently the same apparatus was employed as that ordinarily used in water analysis, and after the large amount of soda required to render 500 c.c. of the jar-water alkaline, was repeatedly distilled with *aqua puriss.* until it gave no further reaction of ammonia, the 500 c.c. was added, and the distillations continued. This determination corresponded to 5.82 mgrms. H_3N in the total jar-water, and is just one-half the amount obtained after reduction. Since the absence of nitrous acid was proven, this result goes far to demonstrate that the ammonia derived from the jar-water, existed there as ammonium nitrate.

But in the ozone wash-water, instead of being one-half, it is only one-third of the amount found on reduction, a result corresponding with that found in the former trial.

Hydrogen Peroxide.—As determined directly upon one portion of the ozone wash-water, this amounted to 2.7 mgrms.; determined after evaporating 30 c.c. down to 3 c.c., it amounted to 2.4 mgrms. The estimation was made by means of sodium hyposulphite and potassium iodide. These reagents could be safely made use of, since the absence of nitrous acid had been previously proven. It was likewise found that the presence of phosphorus acid had no effect upon the titration. If the larger result obtained on titrating directly be attributed to dissolved ozone, then the amount of ozone in solution would have been 0.44 mgrm. Since, however, the percentage of hydrogen peroxide is diminished by concentration with the aid of heat, the lower result obtained after evaporation, might, perhaps, be more satisfactorily attributed to the loss of hydrogen peroxide than to expulsion of dissolved ozone.

The percentage of hydrogen peroxide as compared with the total weight of the air drawn over is 0.00038; that of nitric acid in the ozone wash-water only 0.00008. The former agrees quite closely with the result obtained in the previous experiment; the latter is about one-half.

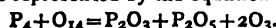
Conclusion.—It will be noted that the chief by-product of the ozonisation of moist air by phosphorus, according to our determinations, is not phosphorous acid, but phosphoric acid. It is generally stated that the former of these two substances is the one principally produced under these circumstances. This may be true in the sense that the phosphorous acid is first formed, and that it gradually is transformed into phosphoric acid under the influence of nascent ozone. This point could not be very readily determined in the course of the experiments made with the apparatus described above.

It is certainly an error to ascribe, as is done in vari-

text-books, the dense white fumes seen in the ozonising chambers, to ammonium nitrite—they are chiefly due to phosphoric anhydride.

The above experiments do not permit us to say that no ammonium nitrite was found during any period of the ozonation, they prove merely that no ammonium nitrite could be detected at its close. If, therefore, this body were produced, it must have become oxidised to ammonium nitrate.

As to the reason of the formation of ozone itself under these circumstances, it may be conjectured along with Lamont* and others, that it is connected with the uneven quantivalencies of the elements taking part in the reaction, which may be represented by the equation—



If this hypothesis be true, then we should anticipate the development of ozone whenever oxidation of a perissad occurred at temperatures compatible with the stability of the ozone molecule. Even at the temperature of combustion of hydrogen, this is supposed by C. Than† to be the case. He explains in this manner the presence of the ozone, which he states he has detected in the combustion of hydrogenous substances generally, and its absence in the combustion of carbon.

In entering into new combinations, the oxygen molecules must undergo temporary resolution into their constituent atoms. These, while *en route* to take up new positions in other combinations, and animated by their atomic energy, or energy of the nascent state, may either oxidise the oxygen molecule, or the nitrogen, or the molecule of water. In the first case, ozone would be produced; in the second, regarding water as the basic body and NNO as the nitryl, there might be formed, as Hunt has indicated, ammonium nitrate; in the third, hydrogen peroxide.

It gives me much pleasure to acknowledge the co-operation of my assistant, Dr. Edgar Everhart, in the performance of these experiments.—*Journal of the American Chemical Society.*

NEW PROCESS FOR THE RAPID ESTIMATION OF PURE SUGAR IN RAW AND REFINED COMMERCIAL SUGARS.‡

By P. CASAMAJOR.

PART I.

THE process for the estimation of sugar, which I propose to describe this evening, is based on a happy idea of M. Dumas. It cannot but be a matter of surprise that, although this idea was published several years ago, it has never been taken up and studied by the numerous chemists whose business it is to analyse sugar.

As an introduction to the subject, I will recall to your memories the main points relating to the process of M. Dumas, and, as this process was suggested to its eminent author, while studying that of Payen, we may also say a few words of the latter.

In 1846, Payen published a process for determining the amount of pure sugar in commercial sugars, which seems to conform more closely to the ordinary processes of chemical analysis than is the case with those which have been more generally adopted for testing sugars. This process consists in washing commercial sugars with alcohol previously saturated with pure sugar. The object of this operation was to wash out everything except the pure sugar, and this, after being freed from the sugar-saturated alcohol by washing with absolute alcohol, could be dried and weighed, as is done with other substances submitted

to chemical analysis. Payen found, however, that the quantity of sugar obtained in this way was always less than the actual quantity present. He seems afterwards to have paid very little attention to this process, and apparently did not attach much importance to it.

It was reserved for Dr. Scheibler, of Berlin, in these latter years, to discover that the result obtained by the process of Payen was not the quantity of pure sugar present, but that it represents the *rendement*, or quantity of sugar obtainable in refining. It cannot but be a matter of surprise that Dr. Scheibler did not distinctly claim this discovery, instead of pretending that he had, in a manner, re-discovered the process of Payen, "which had fallen into the domain of history," when, indeed, he had only attached to it cumbersome paraphernalia, which really added nothing to the value of the process.

Payen used two alcoholic solutions saturated with sugar, and finally absolute alcohol, to wash out the last traces of the sugar-saturated solutions. The first solution was obtained by taking alcohol at 85 per cent, and adding to this 5 per cent of strong acetic acid. This mixture was saturated with sugar. I believe that this addition of 5 per cent of acetic acid was to decompose the sucrales, which were a great bugbear to the sugar chemists of those days. This addition of acetic acid fulfils, however, a useful purpose, as it seems to make the mixture better able to remove the impurities of gummy sugar.

This first solution of Payen was the one adopted by M. Dumas, and here is the process which he proposed for analysing commercial sugars:—

If we introduce an alcohometer of Gay-Lussac in the first solution of Payen, we find that it sinks to a point which, if we operate at 15° C., corresponds to 74 per cent. If we take a certain volume of this solution and stir it up in a glass with a sufficient quantity of a sugar to be analysed, we are able to ascertain, by again placing the alcohometer in the solution, that this has taken up something from the commercial sugar, for the alcohometer no longer indicates 74 per cent, but a lower degree, corresponding to a greater density. The process which M. Dumas proposed was this: Take 100 cubic centimetres of the first solution of Payen and 50 grammes of sugar, agitate the mixture in a glass, filter the solution, and observe the alcohometric degree corresponding to 15° C. For every percentage of sugar less than 100 you will find that the solution indicates 1 per cent less than 74.

A simpler way of stating the same thing is to note that the difference between 100 and 74 is 26, and, if the above proposition is correct, we can obtain the percentage of sugar by adding 26 to the alcohometric degree. Thus, if the alcohometer indicates 68, the percentage of sugar is $68 + 26 = 94$.

The only account I have seen of this process is in L'Abbé Moigno's *Saccharimétrie Physique, Chimique et Mélassimétrie*; this is translated in the *American Chemist* of February, 1871, p. 302. This description may also be found *verbatim* in a recent work of M. Maumené on sugar.

The account of the process of M. Dumas, as found in the above paper, states that, for sugars having 87 per cent or more of pure sugar, the results agree very closely with those of the saccharometer, even within 1-10th of 1 per cent. For sugars of lower grade, the results obtained were not satisfactory.

After making a great number of tests with the solution used by M. Dumas, I am in a position to state that it is not possible, even with sugars of high grade, to obtain results at all approaching those of the optical saccharometer, when we operate on such sugars as we have in this market. The sugars to which the process was applied in France may have been beet sugars of about the same grade and similar composition. With our cane sugars, both raw and refined, the differences between the results of the saccharometer and those of the alcohometer were sometimes as high as 3 or even 4 per cent for sugars above 87 per cent. As to beet sugars I regret to say that,

* CHEMICAL NEWS, vol. xxviii., p. 236.

† *Journ. Chem. Soc.* [2], vol. ix., 1871, p. 483; *Journ. Prakt. Chem.* [2], i., 415.

‡ Read before the American Chemical Society, June 5, 1879.

since I undertook these researches, I have not been able to obtain any samples to form an idea of their behaviour with this process of M. Dumas.

At any rate, as nearly one-half of the raw sugars that come to this market stand below 87 per cent, there seemed to be little use in a process which was declared to be inapplicable to sugars of low grade. I found, however, after trying the process several times, that, although the results obtained were mostly unfavourable, it was impossible to dismiss it entirely, for, upon reflecting upon these results, I found that many questions arose which required to be solved, and, on their solution, I based the hope of modifying this process so as to apply it to the analysis of cane sugars of all grades.

In considering commercial sugars, with the view of applying to them the process of M. Dumas, there are several points which experience has shown to be worthy of attention. Several other points, which have taken a great deal of my time, were found to be of no importance whatever in the study of this question.

To understand the points that are of importance in this inquiry, we may consider that a commercial sugar is composed of four classes of substances:—

1st. Pure sugar.

2nd. Water.

3rd. Soluble impurities, or other soluble substances, besides pure sugar.

4th. Sand, earth and other insoluble substances.

We may now examine each class separately.

1st. The behaviour of pure sugar with the saturated alcoholic solution used by M. Dumas is easily understood. This solution, being incapable of dissolving any further quantity of sugar, its density cannot be affected by the addition of pure sugar alone.

2nd. An addition of water lowers the alcoholic degree. When an excess of sugar is present, the alcohol, diluted with water, is able to dissolve an additional quantity of sugar, which still further increases its density. To ascertain the quantities of pure sugar which alcohol of various strengths will hold, I took several saturated solutions and introduced them in the tube of the saccharometer. The results of the observations thus obtained are given in Table No. 1. In the first column of this table is the percentage of alcohol before being saturated with sugar; in the second column is the indication of the alcoholometer for the same alcohol, after this has been saturated; in the third column is the direct reading of the optical saccharometer (Ventzke's), when the saturated solution is placed in its tube. Finally, in a fourth column, I have placed the number of grammes of sugar in 100 c.c. of the saturated alcohol. The quantities in this fourth column are calculated from those in the third, by taking 0.26 gr. of sugar as corresponding to one degree of the saccharometer scale.

TABLE NO. 1.

Degree of the Alcoholometer before saturation.	Ditto after saturation.	Degree of the saccharometer.	Grms. of sugar in 100 c.c.
92.0	91.0	2.2	0.572
87.0	82.3	—	—
85.0	79.5	12.2	3.17
84.0	78.0	14.6	3.80
82.4	74.0	19.3	5.02
80.0	67.62	26.3	6.83
75.0	48.55	30.0	13.01
50.0	sp. gr. 1.14	192.0	49.9

We may notice, in the above table, that if we compare alcohol of 85 per cent with alcohol of 80 and 75 per cent, that a fall of 5 per cent from 85 to 80 answers to a difference of 11.88 per cent in the corresponding saturated solutions, while a fall of 5 per cent from 80 to 75 corresponds to a difference of 19.07 per cent in their saturated solutions.

These facts are of prime importance in this inquiry, and we will have occasion in the sequel to draw from them consequences worthy of attention.

3rd. *Soluble Impurities.*—To study the effect of soluble impurities on alcohol saturated with sugar, I prepared quite a large quantity of inverted sugar by heating with hydrochloric acid. The acid was afterwards thrown down as chloride of lead, and a small quantity of lead which remained in solution was precipitated with carbonate of soda, but not enough of this reagent was used to make the solution alkaline. This solution was evaporated over a water-bath, and the result was, on cooling, a very stiff gummy substance of light yellow colour. The presence of a slight excess of sodic chloride did not interfere with the usefulness of the product, as it merely acted as the representative of the soluble impurities of commercial sugar, and these always contain soluble salts.

The effect of dissolving inverted sugar in alcohol saturated with sugar was to lower the alcohometric degree, and this lowering was progressively greater as the quantity of inverted sugar was increased. Thus, 5 grms. of inverted sugar lowered the alcohometric degree 11.63 per cent, and 10 grms. lowered it 24.5 per cent, while, if the decrease had been strictly proportional to the quantity of inverted sugar, 10 grms. should have given $11.63 \times 2 = 23.26$ per cent. We may note, however, that the lowering of the alcoholic degree by inverted sugar, which here stands as the representative of the soluble impurities, although not exactly proportional to the quantity added, is more nearly so than is the case with equivalent quantities of water, as we may see in Table No. 1.

4th. *Insoluble Impurities.*—These have no effect whatever on the density of an alcoholic solution saturated with sugar. With clean, dry sand, as well as with pure, dry sugar, no effect takes place in the density of the alcoholic solution. When insoluble substances occur in appreciable quantities, they should be deducted from the indication of the alcoholometer.

It must be understood that in the experiments which are related above with water and inverted sugar, an excess of pure, dry sugar was always present. Otherwise the results obtained would not have been applicable to the analysis of sugars.

Although four classes of substances have been mentioned as worthy of consideration, only water and the soluble impurities have any influence in the results obtained by the process of M. Dumas. If a certain weight of water lowered the alcohometric degree to exactly the same extent as an equal weight of soluble impurities, the process proposed by M. Dumas could, by taking a less weight of sugar for 100 c.c. of solution, be made to yield accurate results; but this does not happen, and if we have, for instance, a sugar containing 94 per cent of pure sugar, the other constituents may be either 6 per cent of water or 6 per cent of impurities, or any sum of impurities and water amounting altogether to 6 per cent. In each of these cases we may obtain a different result. On the other hand, as the addition of a certain quantity of water gives very different results, according to the strength of the alcohol solution, we can conceive that, by a series of experiments, we may succeed in obtaining a solution of such strength that, within the limits which correspond to commercial sugars, the results in lowering the alcoholic degree may be practically the same for a certain quantity of water as for the same quantity of soluble impurities.

Following up this idea, I made a great many experiments, to ascertain whether it was not possible, by raising or lowering the alcohometric degree of a saturated sugar solution, to obtain exactly the strength that would give the required result. In this connection, I may mention, that at an early stage of my experiments, I discarded the use of acetic acid, and I operated with mixtures of alcohol and water only. I found that the solution used by M. Dumas could be replaced by alcohol of 82.4 per cent, saturated with sugar. I tried other solutions, some stronger and others weaker, and I was not long in finding that the bad results obtained with either the solution of M. Dumas or with alcohol of 82.4, saturated with sugar

were much aggravated when the solutions had less alcoholic strength. By a series of experiments, I finally came to the conclusion that a solution of 87 per cent alcohol would give the most satisfactory results. Alcohol of this strength, when saturated with sugar, stands at 82.3 by the alcohometer. This solution answered very well with sugars of quite high grade, but when I came to apply it to gummy sugars, the impurities would not dissolve, and the results obtained were always a great deal too high. I

sought to remedy the evil by taking alcohol at 94 per cent, and lowering the degree with acetic acid instead of water, because, as I have said, acetic acid has a favourable action in dissolving gummy impurities, but I did not succeed in effecting this object, and, much to my disappointment, I had to give up, for a time at least, the project of applying the idea of M. Dumas to the analysis of cane sugars.

(To be continued.)

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

JULY, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

	Appearance in 2 foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	As- Sulphuric hydride.	Hardness on Clark's Scale.								
		Grains.	Grains.								Grains.	Grains.	Grains.	Grains.	Grains.	Grains.	Grains.	Before Boiling.	After Boiling.
<i>Thames Water Companies.</i>																			
Grand Junction	Clear	0.000	0.011	0.105	0.139	21.74	8.500	0.514	1.080	1.16	14.2	3.0							
West Middlesex	Clear	0.000	0.011	0.095	0.135	21.00	7.050	0.504	1.080	1.40	13.3	3.0							
Southwark and Vauxhall	Clear	0.000	0.008	0.120	0.133	19.80	7.160	0.468	1.000	1.43	14.2	3.8							
Chelsea	Clear	0.000	0.010	0.120	0.105	19.00	7.400	0.468	1.080	1.84	14.0	3.0							
Lambeth	Clear	0.000	0.006	0.094	0.105	20.80	8.170	0.720	1.080	1.40	13.2	4.8							
<i>Other Companies.</i>																			
Kent	Clear	0.000	0.001	0.300	0.001	27.90	8.560	0.750	1.512	2.06	18.2	5.1							
New River	Clear	0.000	0.003	0.135	0.056	19.80	7.530	0.470	1.080	1.43	14.0	4.0							
East London	Clear	0.000	0.007	0.105	0.066	21.80	7.280	0.504	1.220	1.36	14.0	3.6							

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 6, 1879.

Sulphurea of Dimethyl-para-phenylen-diamin.—A. Baur.—This compound has the formula $C_{17}H_{22}N_4S$.

Action of Oxalic Ethyl-ether upon Dimethyl-para-phenylen-diamin.—R. Sendtner.—The compounds here examined are dimethyl-para-phenylen-diamin-oxamic ethyl-ether, a body of a basic nature, insoluble in water, and but sparingly soluble in ether and cold alcohol, and dimethyl-para-phenylen-diamin-oxamic acid, mono-para-amido-dimethyl-phenyl-oxamid, and the corresponding di substance.

Phenyl-glyceric Acid or Styeric Acid.—R. Anschütz and L. P. Kinnicutt.—An account of some of the salts of this acid, and of its transformation into dibenzoyl-phenyl-glyceric ethyl-ether.

Desoxalic Acid and its Decomposition into Tartaric Acid.—H. Brunner.—The author has ascertained that the formula of desoxalic acid is $C_6H_8O_9$, and that it forms salts with three or four equivalents of metal. He describes the action of ammonia upon the ether, and the formation of tartaric acid from non-crystalline desoxalic ether.

On Ytterbia.—L. F. Nilson.—The author considers that Marignac had good grounds for assuming that ytterbia is perfectly white, gives no absorption bands, and that its molecular weight = 131. Nilson having taken up

the investigation obtained, after thirteen successive decompositions, an earth of the remarkably low molecular weight 127.6, which he traced to the partial presence of erbia. He hopes to complete the proof that the atomic weight of ytterbia is 132, that it is quadrivalent, and forms a sesquioxide.

On Scandium, a New Earthy Metal.—L. F. Nilson.—Scandium, a metal separated from ytterbia but not as yet obtained in a state of absolute purity, is a white earth whose solutions give no absorption bands. After ignition it is attacked but slowly by dilute nitric acid, but more readily by hydrochloric. The solution of the nitrate is completely precipitated by oxalic acid. The nitrate is completely decomposed at a temperature at which ytterbium nitrate is only partially resolved into a basic salt. The atomic weight, calculated on ScO , must be below 90. The author doubts, however, this composition of the oxide, and thinks the formula ScO_2 more probable. Scandium would then rank between tin and thorium, and with an atomic weight of about 170 it would break the vacancy now intervening between 118 and 234.

Constituents of Combustible Gases in the Potash Mines at Stassfurt.—H. Precht.—In some parts of these mines a combustible gas is liberated in considerable quantities. If mixed with common air it becomes explosive and may give rise to serious accidents. Its composition is:—

Hydrogen	93.053 vols.
Methyl hydrogen	0.778 "
Nitrogen	5.804 "
Carbonic acid	0.180 "
Oxygen	0.185 "
Carbonic oxide	trace
	100.000 "

The author ascribes the origin of this large proportion of hydrogen to the oxidation of ferrous chloride in the absence of air. $6\text{FeCl}_2 + 3\text{H}_2\text{O} = 2\text{Fe}_2\text{Cl}_6 + \text{FeO}_3 + 6\text{H}$.

Measurements of the Density of Organic Solids.—H. Schröder.—The following uniformities appear from the results:—The author shows that in the silver salts of the fatty acid series, and in the ether-sulpho salts of potassium and barium, that for every CH_2 in their composition the volume increases by 3 steres; and that the elements C, H, O, and N, in the solid state, generally occupy the volume of one stere.

Bernardinite, a Resinoid Mineral from San Bernardino, California.—J. M. Stillman.—The composition of this mineral is:—

Water	3.87
Carbon	64.46
Hydrogen (not as water)	8.75
Oxygen	22.80
Ash.. .. .	0.12

100.00

Croton-aldehyd and its Homologues.—A. Lieben and S. Zeisel.—The authors have succeeded by the action of a solution of sodium acetate upon propion-aldehyd in obtaining a condensation product homologous to croton-aldehyd.

Sulphalizaric Acid.—C. Graebe.—The potassium and sodium salts of this acid are soluble in water; those of the alkaline earths and of lead insoluble or sparingly soluble. The salts in which one atom of hydrogen is replaced by a metal are orange or yellow, and if heated in a dry state they yield alizarin. The behaviour of the free acid is similar. Where two atoms of hydrogen are replaced no alizarin is produced, save in case of ammonia. The compounds of the alkalis are a reddish violet, and those of barium and calcium orange. Where three atoms are replaced the behaviour is similar; the salts are violet and are the most readily soluble.

Sulphin Compounds of the Thio-ureas.—A. Bernthsen and H. Klinger.—The authors have obtained a permanent base by the action of benzyl-chloride upon thio-urea.

Purification of Mercury.—J. W. Brühl.—With reference to the method proposed (*Berichte*, xii., 437) by Lothar Meyer, the author states that he tried it some time ago, but with unsatisfactory results.

Derivatives of Cantharidin and their Relations to the Ortho Series.—J. Piccard.—From an examination of cantharen and cantharic acid the author concludes that the cantharidin derivatives are in the ortho series what the camphor derivatives are in the para series.

Fluor-benzol-sulphonic Acid and on Melting-points of Substituted Benzol-sulphon Compounds.—W. Lenz.—No regularities have been detected in the ortho series and the meta compounds have not been sufficiently studied. But as regards the amides and chlorides of the para series the melting-points rise with the increasing molecular weight. The author gives a caution against certain filter-papers containing lignin, which gave rise to perplexing colour reactions.

On α - and β -nitro-alizarin and β -amido-alizarin.—E. Schunck and H. Roemer.—Not suitable for useful abstraction.

On Anthrol.—C. Liebermann and O. Hörmann.—A description of sodium anthracen monosulphate and the corresponding barium and lead salts; of anthrol, acet-anthrol, anthrol-ethyl-ether, and acetyl-mono-oxy-anthraquinon.

Sulphuretted Dyes Obtained from Dimethyl-phenylen-diamin.—A. Koch.—The author has determined the composition of the blue and the red dyes obtained by the patented process of the Baden Aniline Company (No. 1886).

Diphenyl-sulpho-hydantoin.—A. Lange.—Not suitable for useful abstraction.

Dinitro-hydro-cinnamic Acid and its Derivatives.—S. Gabriel and J. Zimmermann.—Among the derivatives here described are the ethyl-ether, amido-nitro-hydro-cinnamic acid, dibrom-amido-hydro-carbo-styryl, and the corresponding mono compound.

Sulphamin-mesitylenic Acids and the Second Oxy-mesitylenic Acid.—Oscar Jacobsen.—Not suitable for abstraction.

Mercuric Iodide.—H. Kœhler.—Mercuric iodide melts not, as stated in chemical text-books, at 238° , but between 253° to 254° , and forms, not an amber-yellow, but a blood-red liquid.

Determinations of the Vapour-densities of certain Inorganic Bodies.—V. and C. Meyer.—The vapour of phosphorus pentasulphide has the density 7.63 to 7.67, calculation showing 7.67. The vapour-density of indium chloride is 7.87, calculation giving 15.20 if In_2Cl_6 , or 7.60 if InCl_3 .

Vapour-densities of Aqueous Acids Boiling at Constant Temperatures.—A. Calm.—The author shows that the vapours of formic, of hydrobromic, and of hydrochloric acids are mixtures respectively of 3, 6, and 9 mols.

Paratoluylic and Terephthalic Acids.—H. Fischli.—An elaborate paper, not susceptible of useful abstraction.

Recognition of the Fatty Alcohols.—H. Gutknecht.—The boundary for the application of Meyer and Locher's test is reached in the secondary class with amyl iodide, whilst with primary alcohols it succeeds in the octylic series, and might probably extend still higher.

Constitution of Parabanic Acid.—A. Calm.—Parabanic acid may be regarded as an intermediate anhydride of oxalic acid and urea.

Benzoyl-cyanide and Phenyl-glyoxylic Acid.—L. Claissen.

Amides of Phenyl-glyoxylic Acid.—L. Claissen.—These two papers do not admit of useful abstraction.

Gas-analytical Determination of Hydrogen by Absorption.—W. Hempel.—This memoir requires the accompanying engraving.

Diphenyl-phthalid and Phenol-phthalein.—A. Baeyer.—The author shows that phenol-phthalein is a dioxy-substitution product of the so-called phthalophenon, which is not a keton, but a diphenyl-phthalide.

Phthal-alcohol.—J. Hessert.—This hydrocarbon is converted by potassic permanganate into phthalic acid, whilst the action of nitric acid yields merely phthalid.

Further Examination of the Putrescent Products of Albumen.—E. and H. Salkowski.—Among the products obtained are phenyl-propionic acid, phenyl-acetic acid, succinic acid, indol, skatol, and phenol, besides small quantities of bodies which require further investigation.

Behaviour of Phenyl-acetic and Phenyl-propionic Acids in the System.—E. and H. Salkowski.—Phenyl-propionic acid in the organism is completely transformed into benzoic acid, and finally appears as hippuric acid in the urine. Phenyl-acetic acid is not oxidised, but forms a corresponding hippuric acid, which might be most accurately named phenaceturic acid.

Dichloracrylic Acid from Muco-chloric Acid.—W. Z. Bennet and H. B. Hill.—A description of the former acid and of some of its salts.

Disubstituted Acrylic Acids.—H. B. Hill.—Not suitable for abstraction.

Synthesis of Oxy-ketons by the Introduction of Acid Radicles into Phenols.—O. Döbner and W. Wolff.—An account of the formation and properties of dibenzo-hydro-quinon.

Chemiker Zeitung.
No. 30, July 24, 1879.

According to Dr. Lenden, the oxidising power of the foam of the waves of the North Sea, observed by Prestel, of Emden, is probably due to the presence of hydrogen peroxide.

Antiseptic power of Pyrogallic Acid.—Bovet has proved experimentally, that pyrogallic acid prevents alcoholic fermentation, and deodorises offensive matter, swarming with bacteria. Kolbe and Meyer, on the other hand, show that the antizymotic power of pyrogallic acid is trifling compared with that of salicylic acid.

Direct Determination of Silver in Galena on Volhard's Principle.—C. A. M. Balling.—From 2 to 5 grms. of the galena, according to its supposed richness in silver, are very finely ground and intimately mixed in a porcelain mortar with from 3 to 4 times its weight of a flux composed of equal parts of soda and saltpetre, placed in a porcelain crucible, covered, and heated over a burner to thorough fusion, when the mixture is well stirred with a glass rod. It is then let cool and placed in an evaporating dish partly filled with water, in which the melted matter is softened, dissolved out of the crucible into the dish, which is then heated, and the watery solution is filtered into a flask. The residue on the filter, after being well washed, is rinsed back into the dish, very dilute nitric acid is added, and the whole evaporated to dryness. The dry residue is taken up in water acidulated with nitric acid, heated, and filtered into the same flask in which is the aqueous solution. The residue is washed with hot water, the filtrate is allowed to cool in the flask, ferric sulphate or iron alum is added, and the liquid is titrated.—*Oest. Zeitschrift Berg. u. Hütten.*

Reimann's Färber Zeitung,
No. 28, 1879.

It is said that the "vert acide" of Poirrier is a sulph-acid of the base of the so-called malachite green.

Bindschedler and Busch, of Bâle, have introduced into commerce a crystalline violet dye, said to be the hydrochlorate of benzylated rosanilin.

Examination of Albumen.—Cordillot dissolves the dry samples in 40 parts of distilled water, and judges the solutions by their degree of fluidity and the presence or absence of insoluble matter. A fine precipitate capable of passing a strainer, is more to be dreaded than large insoluble particles. Equal measures of the solutions are poured into test-glasses, set in cold water, heated slowly to a boil, which heat is maintained for a quarter of an hour. The bulkier the coagulum the better is the quantity. Some samples do not yield a clot at all if dissolved in more than 15 parts of water. The author considers Russian blood-albumen as preferable to all others.

Moniteur Scientifique, Quesneville.
August, 1879.

Recent Progress of Chemical Industry, according to the Reports of Prof. Hofmann.—Soda, by M. Landolt.—This memoir, which occupies nearly half the issue, is a continuation of the reports on the progress of the chemical arts during the ten years previous to the Vienna Exhibition.

Condensation of Acid Vapours from Alkali Works.—Dr. Angus Smith.—Translated from an English source.

Reversion of Superphosphates.—H. Joulic.—Taken from the *Comptes Rendus*.

The so-called Catalytic Force.—Donato Tommasi.—The author considers that the combination of oxygen and hydrogen by means of platinum sponge may be explained on thermo-dynamical principles. Porous bodies, as is well known, condense a certain quantity of gases,

variable according to circumstances. This condensation is attended with the liberation of heat. If its quantity is sufficient the gases absorbed enter into mutual reaction.

History of the Isomeric Purpurins.—Dr. H. Morton.—This paper has already appeared in the *CHEMICAL NEWS*.

Constitution of Ultramarine.—R. Rickmann.—Already noticed.

Impurities contained in Glacial Acetic Acid.—M. Bardy.—The actual acid present in the 57 specimens examined varied from 87 to 99.5 per cent. The author finds that the oil of turpentine may serve for determining with exactness the acid present. For this purpose he takes 10 c.c. of the sample, and carefully drops into it oil of turpentine from a burette graduated into tenths of a c.c. until the last drop added dissolves after slight agitation without producing a permanent turbidity. The quantity of oil which may thus be added increases with the quantity of pure acid. In samples above 99.5 per cent in strength the oil dissolves in any proportion. To obtain comparable results the samples operated upon should be at one and the same temperature, 15° being the most suitable.

In practice it is sufficient to add to a known volume of the acid eight to ten times its volume of the oil and to stir two or three times. If the mixture remains clear the strength of the acid is at least 97 to 98 per cent. Otherwise it should be rejected.

M. Antony Guyard announces a new work on the atomic theory and on his novel theory, to which he gives the name of "tomic."

Les Mondes, Revue Hebdomadaire des Sciences.
No. 13, July 24, 1879.

Treatment of Rabies.—It is recommended to brush the inside of the animal's mouth and tongue first with a liquid not named, then with alcohol at 86 per cent in which Caripagero resin had been dissolved, and lastly with a litre of water containing 100 grms. of liquid ammonia.

MISCELLANEOUS.

Mineralogical Society of Great Britain and Ireland.—The Fourth Annual General Meeting, to receive the Report of the Council, and elect Officers for the ensuing year, and for general business, will be held at the Freemasons' Hall, Surrey Street, Sheffield (by kind permission of the authorities), on Friday, August 22, 1879, at 3 p.m. The following papers will be read:—"On the Production of Different Secondary Forms of Crystalline Minerals," by H. C. Sorby, F.R.S.; "New Scottish Minerals," by Prof. M. F. Heddle; "On some Cornish Serpentine Rocks," by J. H. Collins, F.G.S.

TO CORRESPONDENTS.

G. and F. Malthorp.—No simple means known.
Studio.—It is a legal question.

PATENTS.—Mr. Vaughan, F.C.S., British, Foreign, and Colonial PATENT AGENT. Special attention given to Inventions relating to Chemistry, Mining, and Metallurgy. "Guide to Inventors" Free by Post.—Offices, 67, Chancery Lane London, W.C. and 8, Houndgate Darlington.

WILLOUGHBY BROS., CENTRAL FOUNDRY,
PLYMOUTH, Makers of Plant for Sulphuric Acid and Superphosphate Works, also for Tar and Ammonia Distilling.

NORRINGTON'S PATENT,

For the prevention of escape of gases during the charging of kilns and nitre pots. The success of this invention being assured, we are prepared to Contract for fixing this Patent Apparatus to Pyrites Furnaces throughout the Kingdom.

Estimates and Plans furnished on application.

THE CHEMICAL NEWS.

VOL. XL. No. 1030.

BRITISH ASSOCIATION FOR THE ADVANCEMENT OF SCIENCE.

SHEFFIELD MEETING, AUGUST 20, 1879.

INAUGURAL ADDRESS OF THE PRESIDENT,

PROFESSOR G. J. ALLMAN,
M.D., LL.D., F.R.S.S.L. and E., M.R.I.A., Pres. L.S.

It is no easy thing to find material suited to an occasion like the present. For, on the one hand, there is risk that a presidential address may be too special for an audience necessarily large and general, while, on the other hand, it may treat too much of generalities to take hold of the sympathies and command the attention of the hearers.

It may be supposed that my subject should have been suggested by the great manufacturing industries of the town which has brought us together, but I felt convinced that a worker in only the biological sciences could not do justice to the workers in so very different a field.

I am not therefore going to discourse to you of any of those great industries which make civilised society what it is,—of those practical applications of scientific truth which within the last half century have become developed with such marvellous rapidity, and which have already become interwoven with our everyday life, as the warp of the weaver is interwoven with the woof. Such subjects must be left to other occupiers of this chair, from whom they may receive that justice which I could not pretend to give them; and I believe I shall act most wisely by keeping to a field with which my own studies have been more directly connected.

I know that there are many here present from whom I have no right to expect that previous knowledge which would justify me in dispensing with such an amount of elementary treatment as can alone bring my subject intelligibly before them, and my fellow-members of the British Association who have the advantage of being no novices in that department of biology with which I propose to occupy you, will pardon me if I address myself mainly to those for whom the field of research on which we are about to enter has now been opened for the first time.

I have chosen, then, as the matter of my address to you to-night, a subject in whose study there has during the last few years prevailed an unwonted amount of activity, resulting in the discovery of many remarkable facts, and the justification of many significant generalisations. I propose, in short, to give you in as untechnical a form as possible some account of the most generalised expression of living matter, and of the results of the more recent researches into its nature and phenomena.

More than forty years have now passed away since the French naturalist Dujardin drew attention to the fact that the bodies of some of the lowest members of the animal kingdom consist of a structureless, semifluid, contractile substance, to which he gave the name of Sarcodæ. A similar substance occurring in the cells of plants was afterwards studied by Hugo von Mohl, and named by him Protoplast. It remained for Max Schultze to demonstrate that the sarcodæ of animals and the protoplasts of plants were identical.

The conclusions of Max Schultze have been in all respects confirmed by subsequent research, and it has

further been rendered certain that this same protoplast lies at the base of all the phenomena of life, whether in the animal or the vegetable kingdom. Thus has arisen the most important and significant generalisation in the whole domain of biological science.

Within the last few years protoplast has again been made a subject of special study, unexpected and often startling facts have been brought to light, and a voluminous literature has gathered round this new centre of research. I believe, therefore, that I cannot do better than call your attention to some of the more important results of these inquiries, and endeavour to give you some knowledge of the properties of protoplast, and of the part it plays in the two great kingdoms of organic nature.

As has just been said, protoplast lies at the base of every vital phenomenon. It is, as Huxley has well expressed it, "the physical basis of life." Wherever there is life, from its lowest to its highest manifestations, there is protoplast; wherever there is protoplast, there too is life. Thus co-extensive with the whole of organic nature—every vital act being referable to some mode or property of protoplast—it becomes to the biologist what the ether is to the physicist; only that instead of being a hypothetical conception, accepted as a reality from its adequacy in the explanation of phenomena, it is a tangible and visible reality, which the chemist may analyse in his laboratory, the biologist scrutinise beneath his microscope and his dissecting needle.

The chemical composition of protoplast is very complex, and has not been exactly determined. It may, however, be stated that protoplast is essentially a combination of albuminoid bodies, and that its principal elements are, therefore, oxygen, carbon, hydrogen, and nitrogen. In its typical state it presents the condition of a semifluid substance—a tenacious, glairy liquid, with a consistence somewhat like that of the white of an unboiled egg.* While we watch it beneath the microscope movements are set up in it; waves traverse its surface, or it may be seen to flow away in streams, either broad and attaining but a slight distance from the main mass, or else stretching away far from their source, as narrow liquid threads, which may continue simple, or may divide into branches, each following its own independent course; or the streams may flow one into the other, as streamlets would flow into rivulets, and rivulets into rivers, and this not only where gravity would carry them, but in a direction diametrically opposed to gravitation; now we see it spreading itself out on all sides into a thin liquid stratum, and again drawing itself together within the narrow limits which had at first confined it, and all this without any obvious impulse from without which would send the ripples over its surface or set the streams flowing from its margin. Though it is certain that all these phenomena are in response to some stimulus exerted on it by the outer world, they are such as we never meet with in a simply physical fluid—they are spontaneous movements resulting from its proper irritability, from its essential constitution as living matter.

Examine it closer, bring to bear on it the highest powers of your microscope—you will probably find disseminated through it countless multitudes of exceedingly minute granules; but you may also find it absolutely homogeneous, and, whether containing granules or not, it is certain that you will find nothing to which the term *organisation* can be applied. You have before you a glairy, tenacious fluid, which, if not absolutely homogeneous, is yet totally destitute of structure.

And yet no one who contemplates this spontaneously moving matter can deny that it is alive. Liquid as it is,

* In speaking of protoplast as a liquid, it must be borne in mind that this expression refers only to its physical consistence—a condition depending mainly on the amount of water with which it is combined, and subject to considerable variation, from the solid form in which we find it in the dormant embryo of seeds, to the thin watery state in which it occurs in the leaves of *Valisneria*. Its distinguishing properties are totally different from those of a purely physical liquid, and are subject to an entirely different set of laws.

it is a living liquid; organless and structureless as it is, it manifests the essential phenomena of life.

The picture which I have thus endeavoured to trace for you in a few leading outlines is that of protoplasm in its most generalised aspect. Such generalisations, however, are in themselves unable to satisfy the conditions demanded by an exact scientific inquiry, and I propose now, before passing to the further consideration of the place and purport of protoplasm in nature, to bring before you some definite examples of protoplasm, such as are actually met with in the organic world.

A quantity of a peculiar slimy matter was dredged in the North Atlantic by the naturalists of the exploring ship *Porcupine* from a depth of from 5000 to 25,000 feet. It is described as exhibiting, when examined on the spot, spontaneous movements, and as being obviously endowed with life. Specimens of this, preserved in spirits, were examined by Prof. Huxley, and declared by him to consist of protoplasm, vast masses of which must thus in a living state extend over wide areas of sea bottom. To this wonderful slime Huxley gave the name of *Bathybius Haeckelii*.

Bathybius has since been subjected to an exhaustive examination by Prof. Haeckel, who believes that he is able to confirm in all points the conclusions of Huxley, and arrives at the conviction that the bottom of the open ocean, at depths below 5000 feet, is covered with an enormous mass of living protoplasm, which lingers there in the simplest and most primitive condition, having as yet acquired no definite form. He suggests that it may have originated by spontaneous generation, but leaves this question for future investigators to decide.

The reality of *Bathybius*, however, has not been universally accepted. In the more recent investigations of the *Challenger* the explorers have failed in their attempts to bring further evidence of the existence of masses of amorphous protoplasm spreading over the bed of the ocean. They have met with no trace of *Bathybius* in any of the regions explored by them, and they believe that they are justified in the conclusion that the matter found in the dredgings of the *Porcupine* and preserved in spirits for further examination was only an inorganic precipitate due to the action of the alcohol.

It is not easy to believe, however, that the very elaborate investigations of Huxley and Haeckel can be thus disposed of. These, moreover, have received strong confirmation from the still more recent observations of the Arctic voyager, Bessels, who was one of the explorers of the ill-fated *Polaris*, and who states that he dredged from the Greenland seas masses of living undifferentiated protoplasm. Bessels assigns to these the name of *Protobathybius*, but they are apparently indistinguishable from the *Bathybius* of the *Porcupine*. Further arguments against the reality of *Bathybius* will therefore be needed before a doctrine founded on observations so carefully conducted shall be relegated to the region of confused hypotheses.

Assuming, then, that *Bathybius*, however much its supposed wide distribution may have been limited by more recent researches, has a real existence, it presents us with a condition of living matter the most rudimental it is possible to conceive. No law of morphology has as yet exerted itself in this formless slime. Even the simplest individualisation is absent. We have a living mass, but we know not where to draw its boundary lines; it is living matter, but we can scarcely call it a living being.

The President made further allusion to the researches of Haeckel and others on protoplasm, passing to the consideration of the higher stages in the development of protoplasmic beings. On referring to the characteristic spectrum which the colouring matter of chlorophyll presents under the spectroscopic, he said that for our knowledge of its optical properties we are mainly indebted to the elaborate investigations of Dr. Sorby, which have contributed largely to the advancement of an important department of physical science.

To the presence of chlorophyll is due one of the most striking aspects of external nature—the green colour of the vegetation which clothes the surface of the earth; and with its formation is introduced a function of fundamental importance in the economy of plants, for it is on the cells which contain this substance that devolves the faculty of decomposing carbonic acid. On this depends the assimilation of plants, a process which becomes manifest externally by the exhalation of oxygen. And it is under the influence of light on the chlorophyll-containing cells that this evolution of oxygen is brought about. The recent observations of Draper and of Pfeffer have shown that in this action the solar spectrum is not equally effective in all its parts; that the yellow and least refrangible rays are those which act with most intensity; that the violet and other highly refrangible rays of the visible spectrum take but a very subordinate part in assimilation; and that the invisible rays which lie beyond the violet are totally inoperative.

The President also referred to the fact that the action of chlorophyll in bringing about the decomposition of carbonic acid is not, as was recently believed, absolutely confined to plants. Green *Hydra*, and certain green planariæ and other worms, chlorophyll is differentiated in their protoplasm, and probably always acts here under the influence of light exactly as in plants.

It has, indeed, been proved by some recent researches of Mr. Geddes, that the green planariæ when placed in water and exposed to the sunlight give out bubbles of gas which contain from 44 to 55 per cent of oxygen. Mr. Geddes has further shown that these animals contain granules of starch in their tissues, and in this fact we have another striking point of resemblance between them and plants.

A similar approximation of the two organic kingdoms has been shown by the beautiful researches of Mr. Darwin and others. Indeed, all recent research has been bringing out in a more and more decisive manner the fact that there is no dualism in life,—that the life of the animal and the life of the plant are, like their protoplasm, in all essential points identical.

But there is, perhaps, nothing which shows more strikingly the identity of the protoplasm in plants and animals, and the absence of any deep-pervading difference between the life of the animal and that of the plant, than the fact that plants may be placed, just like animals, under the influence of anæsthetics.

When the vapour of chloroform or of ether is inhaled by the human subject, it passes into the lungs, where it is absorbed by the blood, and thence carried by the circulation to all the tissues of the body. The first to be affected by it is the delicate nervous element of the brain, and loss of consciousness is the result. If the action of the anæsthetic be continued, all the other tissues are in their turn attacked by it and their irritability arrested. A set of phenomena entirely parallel to these may be presented by plants.

We owe to Claude Bernard a series of interesting and most instructive experiments on the action of ether and chloroform on plants. He exposed to the vapour of ether a healthy and vigorous sensitive plant, by confining it under a bell-glass into which he introduced a sponge filled with ether. At the end of half an hour the plant was in a state of anæsthesia, all its leaflets remained fully extended, but they showed no tendency to shrink when touched. It was then withdrawn from the influence of the ether, when it gradually recovered its irritability, and finally responded, as before, to the touch.

It is obvious that the irritability of the protoplasm was here arrested by the anæsthetic, so that the plant became unable to give a response to the action of an external stimulus.

It is not, however, the irritability of the protoplasm of only the motor elements of plants that anæsthetics are capable of arresting. These may act also on the protoplasm of those cells whose function lies in chemical syn-

thesis, such as is manifested in the phenomena of the germination of the seed and in nutrition generally, and Claude Bernard has shown that germination is suspended by the action of ether or chloroform.

Seeds of cress, a plant whose germination is very rapid, were placed in conditions favourable to a speedy germination, and while thus placed were exposed to the vapour of ether. The germination, which would otherwise have shown itself by the next day, was arrested. For five or six days the seeds were kept under the influence of the ether, and showed during this time no disposition to germinate. They were not killed, however, they only slept, for on the substitution of common air for the etherised air with which they had been surrounded, germination at once set in and proceeded with activity.

Experiments were also made on that function of plants by which they absorb carbonic acid and exhale oxygen, and which, as we have already seen, is carried on through the agency of the green protoplasm or chlorophyll, under the influence of light—a function which is commonly, but erroneously, called the respiration of plants.

Aquatic plants afford the most convenient subjects for such experiments. If one of these be placed in a jar of water holding ether or chloroform in solution, and a bell-glass be placed over the submerged plant, we shall find that the plant no longer absorbs carbonic acid or emits oxygen. It remains, however, quite green and healthy. In order to awaken the plant, it is only necessary to place it in non-etherised water, when it will begin once more to absorb carbonic acid, and exhale oxygen under the influence of sunlight.

The same great physiologist has also investigated the action of anæsthetics on fermentation. It is well known that alcoholic fermentation is due to the presence of a minute fungus, the yeast fungus, the living protoplasm of whose cells has the property of separating solutions of sugar into alcohol, which remains in the liquid, and carbonic acid, which escapes into the air.

Now, if the yeast plant be placed along with sugar in etherised water it will no longer act as a ferment. It is anæsthesiated, and cannot respond to the stimulus which, under ordinary circumstances, it would find in the presence of the sugar. If, now, it be placed on a filter, and the ether washed completely away, it will, on restoration to a saccharine liquid, soon resume its duty of separating the sugar into alcohol and carbonic acid.

Claude Bernard has further called attention to a very significant fact which is observable in this experiment. While the proper alcoholic fermentation is entirely arrested by the etherisation of the yeast plant, there still goes on in the saccharine solution a curious chemical change, the cane sugar of the solution being converted into grape sugar, a substance identical in its chemical composition with the cane sugar, but different in its molecular constitution. Now it is well known from the researches of Berthelot that this conversion of cane sugar into grape sugar is due to a peculiar inversive ferment, which, while it accompanies the living yeast plant, is itself soluble and destitute of life. Indeed it has been shown that in its natural conditions the yeast fungus is unable of itself to assimilate cane sugar, and that in order that this may be brought into a state fitted for the nutrition of the fungus, it must be first digested and converted into grape sugar, exactly as happens in our own digestive organs. To quote Claude Bernard's graphic account:—

"The fungus ferment has thus beside it in the same yeast a sort of servant given by nature to effect this digestion. The servant is the unorganised inversive ferment. This ferment is soluble, and as it is not a plant, but an unorganised body destitute of sensibility, it has not gone to sleep under the action of the ether, and thus continues to fulfil its task."

In the experiment already recorded on the germination of seeds the interest is by no means confined to that which attaches itself to the arrest of the organising functions of the seed, those namely which manifest themselves

in the development of the radicle and plumule and other organs of the young plant. Another phenomenon of great significance becomes at the same time apparent—the anæsthetic exerts no action on the concomitant chemical phenomena which in germinating seeds show themselves in the transformation of starch into sugar under the influence of diastase (a soluble and non-living ferment which also exists in the seed), and the absorption of oxygen with the exhalation of carbonic acid. These go on as usual, the anæsthesiated seed continuing to respire, as proved by the accumulation of carbonic acid in the surrounding air. The presence of the carbonic acid was rendered evident by placing in the same vessel with the seeds which were the object of the experiment a solution of barytes, when the carbonate became precipitated from the solution in quantity equal to that produced in a similar experiment with seeds germinating in unetherised air.

So, also, in the experiment which proves the faculty possessed by the chlorophyllian cells of absorbing carbonic acid and exhaling oxygen under the influence of light may be arrested by anæsthetics, it could be seen that the plant, while in a state of anæsthesia, continued to respire in the manner of animals; that is, it continued to absorb oxygen and exhale carbonic acid. This is the true respiratory function which was previously masked by the predominant function of assimilation, which devolves on the green cells of plants, and which manifests itself under the influence of light in the absorption of carbonic acid and the exhalation of oxygen.

It must not, however, be supposed that the respiration of plants is entirely independent of life. The conditions which bring the oxygen of the air and the combustible matter of the respiring plant into such relations as may allow them to act on one another are still under its control, and we must conclude that in Claude Bernard's experiment the anæsthesia had not been carried so far as to arrest such properties of the living tissues as are needed for this.

The quite recent researches of Schützenberger, who has investigated the process of respiration as it takes place in the cell of the yeast fungus, have shown that vitality is a factor in this process. He has shown that fresh yeast, placed in water, breathes like an aquatic animal, disengaging carbonic acid, and causing the oxygen contained in the water to disappear. That this phenomenon is a function of the living cell is proved by the fact that, if the yeast be first heated to 60° C. and then placed in the oxygenated water, the quantity of oxygen in the water remains unchanged; in other words, the yeast ceases to breathe.

Schützenberger has further shown that light exerts no influence on the respiration of the yeast cell—that the absorption of oxygen by the cell takes place in the dark exactly as in sunlight. On the other hand, the influence of temperature is well marked. Respiration is almost entirely arrested at temperatures below 10° C., it reaches its maximum at about 40° C., while at 60° C. it again ceases.

All this proves that the respiration of living beings is identical, whether manifested in the plant or in the animal. It is essentially a destructive phenomenon—as much so as the burning of a piece of charcoal in the open air, and, like it, is characterised by the disappearance of oxygen and the formation of carbonic acid.

One of the most valuable results of the recent careful application of the experimental method of research to the life phenomena of plants is thus the complete demolition of the supposed antagonism between respiration in plants and that in animals.

The President remarked that he had endeavoured to give in a few broad outlines a sketch of the nature and properties of one special modification of matter, which will yield to none other in the interest which attaches to its study, and in the importance of the part allocated to it in the economy of nature. Had occasion permitted he might have entered

into many details which he had left untouched; but enough had been said to demonstrate that in protoplasm we find the only form of matter in which life can manifest itself; and that, though the outer conditions of life—heat, air, water, food,—may all be present, protoplasm would still be needed, in order that these conditions may be utilised, in order that the energy of lifeless nature may be converted into that of the countless multitudes of animal and vegetable forms which dwell upon the surface of the earth or people the great depths of its seas.

We are thus led to the conception of an essential unity in the two great kingdoms of organic Nature—a structural unity, in the fact that every living being has protoplasm as the essential matter of every living element of its structure; and a physiological unity, in the universal attribute of irritability which has its seat in this same protoplasm, and is the prime mover of every phenomenon of life.

We have seen how little mere form has to do with the essential properties of protoplasm. This may shape itself into cells, and the cells may combine into organs in ever-increasing complexity, and protoplasm force may thus be intensified, and, by the mechanism of organisation, turned to the best possible account; but we must still go back to protoplasm as a naked formless plasma if we would find—freed from all non-essential complications—the agent to which has been assigned the duty of building up structure and of transforming the energy of lifeless matter into that of living.

To suppose, however, that all protoplasm is identical where no difference cognisable by any means at our disposal can be detected would be an error. Of two particles of protoplasm, between which we may defy all the power of the microscope, all the resources of the laboratory, to detect a difference, one can develop only to a jelly-fish, the other only to a man, and one conclusion alone is here possible—that deep within them there must be a fundamental difference which thus determines their inevitable destiny, but of which we know nothing, and can assert nothing beyond the statement that it must depend on their hidden molecular constitution.

In the molecular condition of protoplasm there is probably as much complexity as in the disposition of organs in the most highly differentiated organisms; and between two masses of protoplasm indistinguishable from one another there may be as much molecular difference as there is between the form and arrangement of organs in the most widely separated animals or plants.

Herein lies the many-sidedness of protoplasm; herein lies its significance as the basis of all morphological expression, as the agent of all physiological work, while in all this there must be an adaptiveness to purpose as great as any claimed for the most complicated organism.

From the facts which have been now brought to your notice there is but one legitimate conclusion—that life is a property of protoplasm. In this assertion there is nothing that need startle us. The essential phenomena of living beings are not so widely separated from the phenomena of lifeless matter as to render it impossible to recognise an analogy between them; for even irritability, the one grand character of all living beings, is not more difficult to be conceived of as a property of matter than the physical phenomena of radial energy.

It is quite true that between lifeless and living matter there is a vast difference, a difference greater far than any which can be found between the most diverse manifestations of lifeless matter. Though the refined synthesis of modern chemistry may have succeeded in forming a few principles which until lately had been deemed the proper product of vitality, the fact still remains that no one has ever yet built up one particle of living matter out of lifeless elements—that every living creature, from the simplest dweller on the confines of organisation up to the highest and most complex organism, has its origin in pre-existent living matter—that the protoplasm of to-day is but the continuation of the protoplasm of other ages, handed

down to us through periods of indefinable and indeterminate time.

Yet with all this, vast as the differences may be, there is nothing which precludes a comparison of the properties of living matter with those of lifeless.

When, however, we say that life is a property of protoplasm, we assert as much as we are justified in doing. Here we stand upon the boundary between life in its proper conception, as a group of phenomena having irritability as their common bond, and that other and higher group of phenomena which we designate as consciousness or thought, and which, however intimately connected with those of life, are yet essentially distinct from them.

When the heart of a recently killed frog is separated from its body and touched with the point of a needle, it begins to beat under the excitation of the stimulus, and we believe ourselves justified in referring the contraction of the cardiac fibres to the irritability of their protoplasm as its proper cause. We see in it a remarkable phenomenon, but one nevertheless in which we can see unmistakable analogies with phenomena purely physical. There is no greater difficulty in conceiving of contractility as a property of protoplasm than there is in conceiving of attraction as a property of the magnet.

When a thought passes through the mind it is associated, as we have now abundant reason for believing, with some change in the protoplasm of the cerebral cells. Are we therefore justified in regarding thought as a property of the protoplasm of these cells, in the sense in which we regard muscular contraction as a property of the protoplasm of muscle? or is it really a property residing in something far different, but which may yet need for its manifestation the activity of cerebral protoplasm?

If we could see any analogy between thought and any one of the admitted phenomena of matter, we should be bound to accept the first of these conclusions as the simplest, and as affording a hypothesis most in accordance with the comprehensiveness of natural laws; but between thought and the physical phenomena of matter there is not only no analogy, but there is no conceivable analogy; and the obvious and continuous path which we have hitherto followed up in our reasonings from the phenomena of lifeless matter through those of living matter here comes suddenly to an end. The chasm between unconscious life and thought is deep and impassable, and no transitional phenomena can be found by which as by a bridge we may span it over; for even from irritability, to which, on a superficial view, consciousness may seem related, it is as absolutely distinct as it is from any of the ordinary phenomena of matter.

It has been argued that because physiological activity must be a property of every living cell, psychical activity must be equally so, and the language of the metaphysician has been carried into biology, and the "cell soul" spoken of as a conception inseparable from that of life.

That psychical phenomena, however, characterised as they essentially are by consciousness, are not necessarily co-extensive with those of life, there cannot be a doubt. How far back in the scale of life consciousness may exist we have as yet no means of determining, nor is it necessary for our argument that we should. Certain it is that many things, to all appearance the result of volition, are capable of being explained as absolutely unconscious acts; and when the swimming swarm-spore of an Alga avoids collision, and by a reversal of the stroke of its cilia backs from an obstacle lying in its course, there is almost certainly in all this nothing but a purely unconscious act. It is but a case in which we find expressed the great law of the adaptation of living beings to the conditions which surround them. The irritability of the protoplasm of the ciliated spore responding to an external stimulus sets in motion a mechanism derived by inheritance from its ancestors, and whose parts are correlated to a common end—the preservation of the individual.

But even admitting that every living cell were a conscious and thinking being, are we therefore justified in

asserting that its consciousness, like its irritability, is a property of the matter of which it is composed? The sole argument on which this view is made to rest is that from analogy. It is argued that because the life phenomena, which are invariably found in the cell, must be regarded as a property of the cell, the phenomena of consciousness by which they are accompanied must be also so regarded. The weak point in the argument is the absence of all analogy between the things compared, and as the conclusion rests solely on the argument from analogy the two must fall to the ground together.

In a lecture* to which I once had the pleasure of listening—a lecture characterised no less by lucid exposition than by the fascinating form in which its facts were presented to the hearers, Professor Huxley argues that no difference, however great, between the phenomena of living matter and those of the lifeless elements of which this matter is composed should militate against our attributing to protoplasm the phenomena of life as properties essentially inherent in it; since we know that the result of a chemical combination of physical elements may exhibit physical properties totally different from those of the elements combined; the physical phenomena presented by water, for example, having no resemblance to those of its combining elements oxygen and hydrogen.

I believe that Professor Huxley intended to apply this argument only to the phenomena of life in the stricter sense of the word. As such it is conclusive. But if it be pushed further, and extended to the phenomena of consciousness, it loses all its force. The analogy, perfectly valid in the former case, here fails. The properties of the chemical compound are like those of its components, still physical properties. They come within the wide category of the universally accepted properties of matter, while those of consciousness belong to a category absolutely distinct—one which presents not a trace of a connection with any of those which physicists have agreed in assigning to matter as its proper characteristics. The argument thus breaks down, for its force depends on analogy alone; and here all analogy vanishes.

That consciousness is never manifested except in the presence of cerebral matter, or of something like it, there cannot be a question; but this is a very different thing from its being a property of such matter in the sense in which polarity is a property of the magnet, or irritability of protoplasm. The generation of the rays which lie invisible beyond the violet in the spectrum of the sun cannot be regarded as a property of the medium which by changing their refrangibility can alone render them apparent.

I know that there is a special charm in those broad generalisations which would refer many very different phenomena to a common source. But in this very charm there is undoubtedly a danger, and we must be all the more careful lest it should exert an influence in arresting the progress of truth, just as at an earlier period traditional beliefs exerted an authority from which the mind but slowly and with difficulty succeeded in emancipating itself.

But have we, it may be asked, made in all this one step forward towards an explanation of the phenomena of consciousness or the discovery of its source? Assuredly not. The power of conceiving of a substance different from that of matter is still beyond the limits of human intelligence, and the physical or objective conditions which are the concomitants of thought are the only ones of which it is possible to know anything, and the only ones whose study is of value.

We are not, however, on that account forced to the conclusion that there is nothing in the universe but matter and force. The simplest physical law is absolutely inconceivable by the highest of the brutes, and no one would be justified in assuming that man had already at-

tained the limit of his powers. Whatever may be that mysterious bond which connects organisation with psychical endowments, the one grand fact—a fact of inestimable importance—stands out clear and freed from all obscurity and doubt, that from the first dawn of intelligence there is with every advance in organisation a corresponding advance in mind. Mind as well as body is thus travelling onwards through higher and still higher phases; the great law of Evolution is shaping the destiny of our race; and though now we may at most but indicate some weak point in the generalisation which would refer consciousness as well as life to a common material source, who can say that in the far-off future there may not yet be evolved other and higher faculties from which light may stream in upon the darkness, and reveal to man the great mystery of Thought?

ADDRESS TO THE MATHEMATICAL AND PHYSICAL SECTION

BY

G. JOHNSTONE STONEY, M.A., F.R.S., Secretary to the Queen's University in Ireland,
President of the Section.

In order that we may understand the present position of Natural Science upon the Earth, we must remember that the universe is in itself one great whole, which includes minds no less than bodies, for thought is as much a phenomenon of what really exists as motion. But though the universe be but one, man with his limited powers is unable to treat it as such, but has to push his investigation of Nature when and where he can. Thus have arisen many sciences which were at first quite isolated. Their separate condition is a mark of the feebleness of our powers of investigation. Their gradual convergence, and especially where any complete contact can be established between them, is the mark that our advancing knowledge is penetrating deeper.

That there are many sciences of Nature, instead of one science of Nature, has its relation, then, to human imperfection. But the coalescence of sciences has commenced, and is steadily taking place; magnetism is no longer isolated from electricity, nor light from heat, nor the power of thinking from the condition of the brain. In all such cases we have got nearer to understanding what is really going on in Nature. There are already many such achievements of science; but, nevertheless, it remains true that human powers of investigation are so narrow, and the use we have made of them up to the present is so short of what we may reasonably look for in the future, that the sciences of Nature are still many, and most of them stand lamentably aloof from one another.

We find, then, in the present passing condition of our knowledge, one group of sciences which investigate the phenomena of consciousness; another distinct group of the biological sciences; and a third, the group of the physical sciences. These are all but parts of the one great investigation of Nature, but for the present they exist almost disconnected, as separate provinces of human inquiry.

When we endeavour to investigate mental phenomena, we are encountered by the complexity and remoteness of the effects which present themselves for examination, and by a deep and unpenetrated obscurity hanging over the interval between them and their causes. In order to make any progress even in the subordinate task of tracing out the relations of these effects to one another, the inquirer finds it necessary to venture upon hypothesis, and in all metaphysical speculation we sadly miss that healthy discipline with which Nature in other branches of science relentlessly refutes our hypotheses if they are wrong. Here, then, is a region in which the plausible may be mistaken for the true; and it is unfortunately certain that it

* "The Physical Basis of Life." See *Essays and Reviews*, by T. H. Huxley.

has sometimes been so mistaken by the ablest human minds.

The biological sciences treat of all the phenomena of living beings except their mental phenomena, which are those which lie most remote from their causes. Here the complication is less, but it is still too great for the human mind to have yet penetrated behind it. We are still occupied with phenomena which lie at a great distance from their real causes. We are accordingly still far beyond the range of the exact sciences. Most of the great discoveries of biological science have been made by estimating the *general drift* of what is taught by a vast number of particular facts. This, it will be observed, is a kind of reasoning that is necessarily more or less inexact, and, as a consequence, it is one which requires wide intellectual training and great experience and tact to handle it with safety. When the investigator has brought these qualifications to his task, astonishing progress has been made in these sciences: without them the reasoning may degrade into being either trivial or loose.

In the rest of the study of Nature we are not embarrassed by the phenomena of life, and many mysteries therefore stand aside out of our path. Here lies the domain of the physical sciences. It is here that the mind of man has best been able to cope with the realities of the universe, and in which its greatest achievements have been effected. It is here that exact reasoning finds a predominant place.

The study of the physical sciences has been remitted by the British Association to its first two sections, chemistry being assigned to Section B, and the rest of the physical sciences to Section A. Accordingly, Section A includes the whole range of mathematics, along with the study of the conditions of rest and motion in that part of matter which is endowed with mass, and of the phenomena of sound, heat, light, and electricity, with the applications of these abstract studies in molecular physics and to astronomy, crystallography, and meteorology.

In meteorology, owing to the complication of the materials that have to be dealt with, we must have frequent recourse to the same kind of reasoning as has been found so effectual in the biological sciences; but in the other physical sciences which I have enumerated exact reasoning prevails, and on this account they are frequently classed together as the exact sciences.

The process of investigation in the exact sciences is fundamentally one in all cases. It has been well described by Mill in the third book of his "Logic." Nevertheless, it is notorious that minds which are well fitted for some branches of physical inquiry find difficulty—sometimes insuperable difficulty—in pursuing others. It is not every eminent mathematician who would have made an equally good chemist, or *vice versa*. This is because there exists a practical distinction separating the investigation of exact science into two well-marked classes when they are viewed, not as they are in themselves, but in their relation to the powers of us human beings. I refer to the distinction between the experimental method or the method of observation on the one hand, and the deductive method, or the method of reasoning on the other. All valid investigations in exact science appeal to what can be directly perceived, and all lead to a conclusion which can be reasoned out from it; but there are some of these investigations in which the main difficulty consists in making the appeal to the senses, and there are others in which the main difficulty lies in the process of reasoning.

To contend with these difficulties successfully requires very different qualities of mind and body. In experimental science the powers principally called into requisition are readiness and closeness of observation, dexterity in manipulation, skill in devising expedients, accuracy in making adjustments, and great patience. It also requires that the investigator should have an accurate memory of what else he has witnessed resembling the phenomenon under observation, that he should be quick to detect every point of agreement and difference that can be perceived,

and be skilful to select those which are significant, and to employ them as materials for provision to guide his further proceedings. But the strain on the reasoning powers is generally less, often of trifling amount. The question is put to Nature, and it is Nature usually that gives the bulk of the answer. The most striking monument of splendid achievements by the experimental method of investigation unaided by the deductive method is to be found in the science of chemistry.

An equally typical instance of the power of the deductive method is the science of mechanics. This science, which has sunk deeper into the secrets of Nature than any other science, and which is the science towards whom all other physical sciences are at present more or less gravitating, is essentially deductive. There is little or no difficulty about its fundamental data. They are facts of Nature so patent to all men, and so indelibly implanted in human conception, that some persons have supposed that we have an intuitive perception of them. But, while the materials from which the mind is to work are thus easily obtained, it has taxed to the utmost the reasoning powers of understandings like Newton's to evolve the few consequences of them which are already known, and the investigator has to call to his assistance every aid to prolonged consecutive thought which mathematicians can devise.

In grappling with the problems of Nature we are seldom allowed the choice of the method of investigation we shall employ. This is commonly settled for us and not by us. Where we cannot advance without further information we must make further observations, *i.e.*, we must employ the experimental method, the appeal *ad experientiam*: where we cannot advance without understanding better what the information we possess really amounts to, we must employ the deductive method.

No reach of intellect applied to the materials in existence before 1860 could have elicited the fact that iron exists upon the sun. This great discovery was made by Professor Kirchhoff, a scientific man who was equally versed in both methods of investigation. On the present occasion it was the experimental method he employed. He applied to the scrutiny of the sun's spectrum four prisms of the most homogeneous glass that could be procured, figured with the greatest accuracy that the eminent artist Steinheil could attain. He expended far more pains on their adjustment for each successive part of the spectrum than any of his predecessors had done, and he was rewarded by a more perfect vision of the sun's glorious spectrum than had met the human eye before. In a collateral inquiry, suggested by an observation made by Foucault, he and Bunsen placed a metallic vapour emitting bright rays in front of a still brighter incandescent body, so that the light from the brighter background had to pass through this vapour, and they found that this vapour now caused dark lines in the spectrum occupying the positions which its own bright lines had before filled. Professor Kirchhoff thereupon added an appliance to his spectroscopic which enabled him to bring a metallic spectrum and the solar spectrum together into the field of view, alongside of one another. On accomplishing this he saw sixty of the brightest of the iron rays as continuations of sixty of the strongest of the dark lines in the sun's spectrum; and, by an elaborate scrutiny, he satisfied himself that the observations had been pushed to a sufficient degree of exactness to make sure that a deviation would have been detected in any one of these sixty cases if it had amounted to as much as one-fourth of the average interval between consecutive lines of the solar spectrum. From this it was obvious that the sixty coincidences are not due to chance, but indicate that there is really iron vapour in the path of the rays. It will be observed that Kirchhoff's great merit and the real difficulty of his work lay in the scientific foresight and the industry which were required to frame hypotheses that were worth testing, to guide the investigation by these hypotheses, to contrive, construct, and adjust adequate apparatus, and to make with it the elaborate observations and the exact observations and maps which were necessary.

But when by these means the new facts had been brought to light, the inference from them that there is iron in the atmosphere of the sun was an easy one. This example will better convey than a definition what are the characteristic features of an experimental inquiry.

On the other hand, no series of observations or experiments, however skilfully arranged, could have enabled anyone to understand the cause of that familiar but truly surprising phenomenon that a top stands upon its peg while it is spinning. But a full explanation of it is within the reach of any student who will train his mind to reason consecutively, and avail himself of the aids to prolonged consecutive thought which mathematicians have contrived. He will then see that the most obvious and familiar mechanical facts involve as necessary consequences all the phenomena which he finds in the schoolboy's top, in the physicist's gyroscope, and in the precision and nutation of the heavens. This, then, is a problem of Nature which falls within the province of the deductive method.

Wherever data are known exactly, these inferences from these data, however remote, may be depended upon as corresponding with what actually occurs in Nature. And if in such cases the mind of man has proved equal to the task of drawing inferences which can effectually grapple with the problems he finds around him in the Universe—which is, alas! as yet but too seldom—then will the deductive method, our plummet, explore depths in the great ocean of existence which our anchors of experiment could not have reached.

The distinction which is here made between deductive and experimental investigations would have no place in a logical system. But it has direct reference to human convenience, and derives its importance from this circumstance. It is obvious, too, that an investigation may partake of both characters—that it may require all the powers of the scientific observer to get at the facts, or even to appreciate them, and all the resources of the mathematician to elicit the consequence of them. For instance, on beginning his electrical studies, the student of Nature must master a mixed experimental and deductive inquiry to get at the elementary fact that free electricity resides either at or outside the surfaces of conductors; and he must engage in a further inquiry, and one only within the reach of a trained mind, to deduce from this the law of the inverse square. And, again, no full appreciation or even intelligent use of the common electrostatic and electrodynamic measures which he meets at the threshold of his electrical studies is within the reach of the mere experimentalist or the mere theorist. And if this treacherous ground lies before the immature student at his entrance, what shall we say of the bogs he struggles into as he advances? We are perpetually meeting with inquiries of this mixed character in electricity and some of the other physical sciences, but they are comparatively rare in either medicines or chemistry, and none that is difficult lies in the path of the beginner. How many students are there who are made to slur over the above and a multitude of similar difficulties, and who are told that they are learning science, when in fact what they are really learning is the pernicious habit of being content to see Nature through a fog or through other men's mental eyes.

In mechanics valuable progress can be made by the mere mathematician, the student of deductive science; and in chemistry similar progress can be made by the mere experimentalist. Of all the physical sciences these are the most purely deductive, and the most purely experimental. What I desire particularly to invite attention to is that the two great methods of investigation may best be acquired in these two sciences, and that for a really sound grasp of the remaining physical sciences, and especially with a view to further advance in physical science, a command of both methods of investigation is essential.

We must bear in mind, too, that either method of investigation may be misapplied, and that this is a risk carefully to be guarded against. The deductive method

when misapplied lands us in speculation; the experimental method becomes empiricism; and it so happens that the sciences of mechanics and chemistry are not only monuments of the power of the two great methods of investigation, but instructive examples of their weakness also. For in chemistry scarce any attempt at prolonged reasoning, carrying us by any lengthened flight to a distance from the experiments, can be relied on. The result has seldom risen to anything better than speculation. And on the other hand, in mechanics, conclusions which depend on experiments only are empirical; that is, they are deficient in accuracy, and their relation to the other phenomena of the science is left in darkness. Here, then, we find in these two sciences not only how strong these two methods of investigation are, but how weak they may become if misapplied.

I do not know whether any of my predecessors in this chair has experienced so much difficulty, or has hesitated so long and so much as I have hesitated in selecting the topic to which he would ask your attention. My first effort was an attempt to delineate the great recent progress of the mathematical and physical sciences, but it was unsatisfactory, partly from my own too scanty powers, and also because the variety, and even disparity, of the numerous sciences somewhat arbitrarily grouped together in Section A gave to the outline too sketchy a character. My next attempt was to make a selection among them, confining myself to those with which I am best acquainted, and endeavouring to direct attention to the problems which at the present time seem most to stand in need of solution. But here I felt unwilling either to bring forward or to withhold views which might be disputed. I then applied myself to the single consideration of what I hoped might prove useful and not inopportune at a time when one university, which I trust will prove a great university, is rising in the north of England, and when another university which has carefully and successfully fostered a high standard of education for thirty years, and which has thereby deserved and won the respect of educated men, has just been sacrificed to ecclesiasticism in the sister isle. In this university I have held the most central office for twenty-two years, and in the discharge of my duty had largely to influence its destiny in respect to almost every educational problem. Parliament in its wisdom has now seen fit to destroy this work, and I have not been without hope that from the experience which has been gained some effect which shall last may yet arise, and that the Queen's University may perhaps at its death bequeath a useful legacy to the University of Victoria. The advancement of Science in the North of England will largely depend for many years on the wisdom of the regulations for scientific training which are adopted at first by the new university; and I have therefore ventured, at this peculiar juncture, to submit to the judgment of my scientific brethren the principles which much thought and many trials extending over several years have led me to believe should guide them in selecting this part of a curriculum.

I have sought to show that it is in the study of mechanics and in the practice of chemistry that the two great methods of investigation may best be acquired. In them they may be studied separately, by steps of graduated difficulty, and with a superabundance of materials; and each of them supplies the necessary cautions with respect to the method which is all powerful in the other. No scientific man is really equipped for the pursuits in which both methods have to be employed till he has separately acquired a grasp of each. For it is only then that he will be armed against the errors which lead so many to mistake empiricism on the one hand, and speculation on the other, for solid science, or to underrate solid science, mistaking it for speculation. Nor is it only in his scientific occupations that he will derive benefit from this training. All exact reasoning, whether in science or in common life, belongs to these great divisions; and in the numberless instances in which we must be satisfied with reasoning

which falls short of being exact, our only safety lies in having by the practice of exact reasoning, both deductive and experimental, attained to that intellectual tact and caution which alone will enable us to handle with safety the sharp and slippery tool. It is thus that a sound judgment with regard to truth may best be acquired by man or woman; and soundness of judgment is the noblest endowment of man's understanding, just as veracity is first among his virtues.

NOTE ON CHARACINE.

By Dr. T. L. PHIPSON, F.C.S., &c.

AMONG the organic substances present in fresh water is a new and interesting product, to which I have given the name Characine, for reasons that will appear presently. It is the substance to which Algae in general—such as *Confervæ*, *Oscillariæ*, *Desmids*, &c., and the plants of the genus *Chara*—owe their peculiar odour, and communicate this odour to the water in which they abound.

I have obtained it, in minute quantities only at present, from *Palmella cruenta*, from *Vaucheria terrestris*, and from several *Oscillariæ* (*Oscillaria autumnalis*, *O. tenuis*, &c.) It is, apparently, more developed in the genus *Chara*, and *C. fastida* will, I believe, yield it in larger quantity than the plants already mentioned. It is also plentifully produced by the dark coloured *Oscillariæ* growing on damp walls and by *Nostocs*.

Characine is a kind of camphor which gives to these plants, and to the waters in which they grow, that peculiar, highly characteristic, marshy odour, which is usually ascribed to products of their decomposition, but which is due entirely to the substance here described, produced by the plant in life and health. It is formed not only in those algae which live entirely in water, but also by those, such as *Palmella* and *Oscillaria*, which flourish in moist places, and are occasionally subjected to desiccation. It is sometimes seen floating in extremely thin films upon the surface of stagnant waters and on that of tanks in which Algae are cultivated. I obtained it first from *Palmella cruenta* whilst studying its curious pink colouring-matter, *Palmelline*, recently described in a note to the Paris Academy (of which remarkable substance I shall soon have more to say).

When a certain quantity of alcohol in which this plant has steeped for some twenty-four hours in a closed tube is diluted with fifteen to twenty times its volume of water, and then (the grains of chlorophyll having been allowed to deposit and the liquid decanted) shaken up with ether, the latter dissolves the characine, and leaves it on evaporation as a white greasy substance, having a strong and characteristic marshy odour; non-saponifiable by potash; lighter than water; gradually volatilising into the air (or disappearing by oxidation) from the surface of water on which it floats, and which thus loses its odour entirely in two or three days; soluble in alcohol and in ether, but nearly insoluble in water. (When heated with water in a closed tube it was transformed into a substance similar to vegetable wax, melting at 83° C., and having the odour of that substance; but this effect might have been due to some impurity). A better method of obtaining characine in a pure state is not to employ alcohol, but to operate as follows:—

The *Palmella* or *Oscillaria* which is to be treated must be previously dried by exposure to the air, at a temperature not exceeding summer heat, for about twenty-four hours. The "dry" substance is then covered with cold water in a capsule, which must itself be covered with a sheet of glass, and in the course of about thirty-six hours more (with *Palmella cruenta*) thin films of characine will be observed floating on the water. The latter is then decanted off into a long tube, together with the films (which are apt to stay behind in the capsule), and shaken

up with ether as before mentioned. In this case the product is perfectly white, quite devoid of crystallisation, more or less unctuous in appearance, whilst that obtained by the use of alcohol has often a yellowish tint, and is probably impure. Up to the present time I have not obtained it in sufficient quantity to ascertain more of its properties.

BLEACHING OF SUGAR SYRUPS BY OZONE.

By ALBERT R. LEEDS.

NOT being able to find in chemical journals any accounts of experiments upon coloured syrups with ozone as a bleaching agent, while there were rumours that many such had been tried, it appeared the readiest way of obtaining information to institute suitable trials, with quantitative determinations of the amounts of ozone employed and of the degrees of change effected.

The material was kindly furnished by Dr. Arne Behr, from syrups manufactured in the refinery of Messrs. Matthieson and Weichers, Jersey City. The first specimen was of syrup which had undergone but one filtration, and was of a brownish yellow colour. In this preliminary experiment the amount of ozone required to effect the bleaching was not determined. At its close, the syrup was of a faint straw colour and of slightly acid reaction.

A second trial was made upon a syrup which had been twice filtered, still retaining a strong yellow tint. 20 c.c. of the syrup were introduced into a Geissler absorption apparatus, and a slow current of oxygen, ozonised to the extent of 24 m.grms. ozone per litre, passed through it for several hours. When about 100 m.grms. ozone had been brought into contact with the syrup it had become almost colourless. To my own litmus-paper it was neutral, although Dr. Behr informs me he detected a very feeble acid reaction.

As determined by Dr. Behr, the filtered syrup when it came from the refinery contained, in 100 parts, 50 parts of dry substances and 40 parts of dry sugar. The alteration in the course of bleaching is seen in the following table:—

Effect of Oxone upon Filtered Syrup.

Dry Substance contains.	Unbleached.	Bleached.
Cane-sugar (by polariscope) ..	79.7 p.c.	80.0 p.c.
Inverted sugar	12.7 "	12.7 "

ACTION OF OZONE UPON THE COLOURING MATTERS OF PLANTS.

By ALBERT R. LEEDS.

IN preceding numbers of the *CHEMICAL NEWS* for this year I have given the results obtained when ozonised oxygen was made to act upon flowers and plants. They were exposed to the action of a slow continuous current of ozonised oxygen, containing 1 m.grm. ozone per litre, during intervals varying from eighteen to thirty-six hours. The results, which were mainly negative, were so unsatisfactory that I repeated the experiments, with the aid of more efficacious apparatus for generating ozone.

One phosphorus-ozonator was employed to supply ozone; a second, without the phosphorus-cakes upon the disks, made a convenient arrangement to contain the flowers. As their stems were placed in vessels filled with water, and the bells were closed by the water contained in the jars, the atmosphere surrounding the flowers was kept moist, which was of importance in conducting the experiments.

I. In the first trial, in which many varieties of flowers were exposed during nineteen hours to the

action of a current of 152 litres of air, containing in all 228 m.grms. of ozone, the bleaching effected was extremely imperfect.

II. 1200 Litres of Air passed over. (Total ozone 1·8 grms.)
At the end of five days—

Lantana, red	Yellow, and somewhat decomposed
Fuchsia, calyx, carnation-red	Yellow
„ petals, rose-pink	Dirty white
Petunia, magenta coloured	Calyx and petals partly bleached
Rose, crimson	Yellow
Agapanthus umbellatus	White at end of petals
Verbena, purple with white centre	Bleached
Pelargonium, scarlet	Bleached
Euphorbia, salmon coloured	Partly bleached
Fuchsia	Calyx and stamens yellow
Verbena, maroon coloured	Dirty white

A piece of calico with a pattern in bright green and black was completely bleached during the same interval, the green colour having disappeared completely, and the stain of the mordant only remaining where the black had been.

III. Ozonised Oxygen. (Exposure three and a half hours.)

Rose, light red	Nearly white; leaves white in spots
Fuchsia	Bleached to light red
Verbena, purple	White with purple spots
„ red	White with red spots
Petunia, purplish red	White in spots
Pelargonium, pink	White at end of petals
Bouvardia	Dirty white

No determination was made on the percentage of ozone during this particular experiment. But the same electrical ozoniser was employed as in other trials, when it had produced 24 m.grms. ozone per litre. A slow current of the gas was kept flowing during the course of the entire experiment.

Conclusion.—The colouring matters of both leaves and flowers of the species experimented upon were partly or wholly destroyed by ozone; but a considerable percentage of ozone is required to produce this result, or if such small amounts as are obtained in the customary methods of ozonising air by phosphorus are employed (1 to 3 m.grms. per litre), a large volume of ozonised air must be used, and a considerable interval elapse, before bleaching is effected.

THE BROMINE DERIVATIVES OF β -NAPHTHOL.

By ALFRED JOHN SMITH, B.Sc.

I HAVE succeeded in obtaining for the first time a mono- and tetra-bromine derivative from β -naphthol. The mono-bromide is formed by adding a mixture of bromine and glacial acetic acid drop by drop to a solution of β -naphthol in glacial acetic acid. On standing, it crystallises out in colourless needles, which can be washed from acetic acid with distilled water, and dried between filter-paper. The tetra-bromide is formed by adding excess of bromine to the solution of β -naphthol in glacial acetic acid; it crystallises out quickly. It is washed from excess of bromine with glacial acetic acid, then crystallised till quite white, washed with water, and dried in the same way as the mono-bromide.

The mono-bromide is decomposed by the action of heat, giving off copious fumes of hydrobromic acid. The evolution of HBr begins at about 130°; it melts at 84° C. On heating gradually up to 250° C. in an oil bath with excess

of caustic potash or soda and a little water, β -naphthol is again formed.

The effect of an alkaline solution of potassium permanganate, at the heat of the water-bath, was tried both with the mono- and tetra-bromide; the excess of permanganate was destroyed with ferrous sulphate, the solution acidified with hydrochloric acid and extracted with ether. The ethereal extract deposited crystals on evaporation, which were submitted to sublimation. From the mono-bromo compound the anhydride of phthalic acid was obtained, and from the tetra-bromo compound the anhydride of mono-bromo-phthalic acid, thus showing that in the mono-bromo compound the bromine and hydroxyl are in the same benzene nucleus, and that in the tetra-bromo compound three bromine atoms and the hydroxyl are in the same benzene nucleus.

A full account of the research will shortly be offered to the Chemical Society. The research was carried on in the laboratory of the University of Zürich, under the supervision of Professor Merz.

CORRESPONDENCE.

ANALYSTS IN JAPAN.

To the Editor of the Chemical News.

SIR,—Referring to the memorial which I presented at your office, I shall feel obliged if you will briefly notice it in your next issue.

A memorial to the Secretary of State for Foreign Affairs, signed by a number of the leading manufacturers and wholesale export drug and chemical merchants—amongst them Messrs. Howards and Sons, Allen and Hanburys, Burgoyne, Burbidges, and Co., Baiss Brothers and Co., Curling and Co., Dakin Brothers, Corbyn, Stacey, and Co., Herring and Co., Horner and Sons, May, Baker, and Co., Whiffin, &c.,—petitioning his lordship that Her Majesty's Government may take such steps as it may deem advisable to induce the Japanese Government to employ competent officers in the analyses of drugs and medicines, &c., at the ports of Japan open to foreigners; since, during the past two years, whole invoices of the purest and best qualities of such goods have been condemned as impure and prohibited for use and sale in that country, thereby causing heavy losses to merchants engaged in business in that country directly or indirectly.

The memorial is accompanied by several certificates of Professor Atfield and others confirming the purity of the goods condemned by the officials in the Government service of Japan.—I am, &c.,

J. HARTLEY.

9, Charterhouse Buildings, E.C.
August 14, 1879.

ACTION OF HEAT ON IODINE.

To the Editor of the Chemical News.

SIR,—Messrs. V. and C. Meyer, in the last Report of the German Chemical Society, give numbers which show a large diminution in the specific gravity of chlorine at the high temperatures they employed, and state that the behaviour of iodine gas was quite analogous to that of chlorine. These experiments must now be known to everyone, but the words with which the Messrs. Meyer conclude their most important series of papers calls, I think, for comment. They say:—

“For the present we will only refer to the fact, that the measurements of high temperatures, and the vapour density estimations based upon them by Deville and Troost, presuppose the unalterability of the density of iodine vapour. The inadmissibility of this premise being

recognised, partly new interpretations will become necessary for the classical works of these experimenters."

—(*Berichte der Deutsch.*, xii., 1431.)

But Deville and Troost, where they employed an iodine thermometer, also checked their result by an air thermometer, and the Messrs. Meyer infer from their experiments that at 1567° C. (their highest temperature) the molecular weights of oxygen and nitrogen still correspond to the formulæ O_2 , N_2 , usually ascribed to them. —(*Berichte der Deutsch.*, xii., 1428.)

Thus Deville and Troost found the boiling-point of sulphur by air thermometer to be 440° C. The specific gravity of iodine gas, accepting this temperature 440°, was found to be 8.70.

The boiling-point of cadmium, as found by air thermometer, was 856°; the iodine pyrometer gave numbers which, accepting 8.716 as specific gravity iodine gas, gave 860° as boiling-point of cadmium.

Finally, the boiling-point of zinc by air thermometer = 1034°; by iodine pyrometer, assuming 8.716 to be the specific gravity of iodine gas, boiling-point of zinc = 1040°. —(*Ann. de Chimie*, lviii., 257.)

From these numbers it would appear that at 1000° C. the specific gravity of iodine gas does not appreciably alter, but on that point we shall no doubt soon have information from Prof. Meyer, my only object in writing being to point out that the accuracy of the beautiful work of Deville and Troost is not wholly dependent on the indications of the iodine pyrometer.—I am, &c.

W. H. DEERING.

1, Parrock Road, Gravesend.
August 18, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 2, July 14, 1879.

Direct Combination of Cyanogen with Hydrogen and with Metals.—M. Berthelot.—The author, having measured the heat of the formation of hydrocyanic acid and of cyanogen from their elements (−14.1 and −38.3), concludes that the synthesis of hydrocyanic acid from cyanogen and hydrogen ought to evolve a considerable quantity of heat. He finds that gaseous hydrocyanic acid may be heated to 550° for three or four hours in a sealed tube without betraying any marks of decomposition or dissociation. The author effected the direct combination of cyanogen and hydrogen by heating the pure dry gases in equal volumes in a sealed tube of hard glass to 500° to 550° for several hours. On opening the tube a loss of about one-seventh of the volume was apparent, due to the formation of a certain quantity of para-cyanogen. Potassa absorbed five-sevenths of the gas, and the residual one-seventh was found on analysis to consist of water almost pure. The volume of this residual hydrogen being sensibly equal to the original condensation (representing the change of a certain quantity of cyanogen into para-cyanogen) it follows that the gas absorbable by potassa is hydrocyanic acid exempt from free cyanogen. At a lower temperature the synthesis is less complete, and at greater heats a portion of nitrogen is set free. At 300° cyanogen combines with zinc, cadmium, iron if brought in contact in a sealed tube.

Organo-metallic Radicles of Tin: Stannbutyles and Stannamyles.—A. Cahours and E. Demarcay.—Not capable of useful abstraction.

Researches on the Effects of the Rheostatic Machine.—G. Planté.—The length of the sparks given by

this machine is sensibly proportional to the number of condensers. With 80 condensers the author obtained sparks of 0.12 metre in length. If produced over a surface formed of a mixture of resin and paraffin they leave branching furrows, a representation of which is appended to the author's memoir.

Vapour of Bihydrosulphate of Ammonia.—M. Isambert.—The author shows that the experiments of MM. Engel and Moitessier (*Comptes Rendus*, lxxviii.) are not conclusive.

Solution of Carbonic Oxide in Acid Cuprous Chloride.—H. Hammel.—The author has determined the quantity of heat disengaged in the reaction of carbonic oxide upon cuprous chloride, dissolved in hydrochloric acid.

Transformation of Tartaric Acid into Glyceric and Pyruvic Acids.—G. Bouchardat.—This change is produced by the action of sulphuric anhydride. The mucic, citric, and malic acids are decomposed in an analogous manner.

Isomerisms of Borneol.—J. de Montgolfier.—The author has determined the optical rotatory power of certain camphols, and of the camphor regenerated from them.

Bihydrochlorate of Terebenthen.—J. de Montgolfier.—The action of sodium upon the bihydrochlorate gives rise to a mixture of carbides of the formula $C_{20}H_{16}$ and $C_{20}H_{20}$.

Certain Derivatives of Indigotin.—E. Giraud.—The indolin of Schützenberger appears to be generated by a body identical, or, at least, closely connected, with the flavindin of Laurent. It is easily obtained by heating indigotin to 180° along with the hydrosulphite of soda and an excess of caustic soda.

No. 3, July 21, 1879.

Various Thermo-chemical Data.—M. Berthelot.—The author determines the formation heat of diamylen in the gaseous state at +15.4 cal. The melting heat of glycerin is 3.91 cal., and its specific heat is $47.8 + 0.14 t$ (t being the temperature).

Phenomenon Analogous to that of Peltier.—E. Bouty.—If water acidulated with sulphuric acid is electrolysed between two thermometers covered with platinum the temperature is higher at the positive than at the negative pole. If the current is reversed a strong fall is observed at the pole which has become negative, but it disappears in a few moments. If the water is acidified with hydrochloric acid the initial phenomenon produced by the reversal of the current is a great rise of temperature at both poles.

Capacity of Voltaic Polarisation.—R. Blondlot.—Not susceptible of useful abstraction.

Action of Magnetism in Movement upon Static Electricity; Inertia of Static Electricity.—G. Lippmann.—The so-called phenomenon of Rowland (*Berlin Berichte*, 1876) is necessarily reversible, and this reversibility is a consequence of the impossibility of perpetual motion. Static electricity possesses a special mechanical inertia which is simply added to that of the electrified body.

Laws of the Variations of Atmospheric Electricity deduced from the Regular Observations made at the Observatory of Moncalieri.—P. F. Denza.—Atmospheric electricity presents daily in Piedmont two maxima following the rising and setting of the sun, at an interval of some hours. These two maxima are separated by a minimum which follows the passage of the sun over the meridian of the place. As regards the annual fluctuation the maximum value of the atmospheric tension falls in February, and the minimum in September. Before and after storms the electrometer almost always marks zero, but during their passage or proximity the tension is very great. Rain and snow increase tension more slightly and are often preceded and followed by electric diminution.

The action of fogs, hoar frost, and of the formation of clouds increases atmospheric electricity, though to a less extent than that of rain and snow. In calm and hot weather the lowest values are observed. South and especially south-easterly winds increase the electricity of the air; north winds have an opposite effect. Rain and snow are accompanied by negative electricity at least as often as by positive. The same proportion holds good for storms and to a less extent for rain and snow. Negative electricity is generally due to storms or rain at a distance, to the formation of clouds, or to a polar aurora. In the normal conditions of the atmosphere electric tension decreases with altitude.

Researches on Explosives: Combustion of Powder.—MM. Nobel and Abel.—The length of this paper renders it unfit for abstraction.

Experimental Researches on the Decomposition of Gun-cotton in Closed Vessels.—MM. Sarrau and Vieille.—The decomposition of gun-cotton gives rise to products which are few in number and very simple, namely, carbonic acid, carbonic oxide, hydrogen, and nitrogen.

Analytical Use of Sulphuretted Hydrogen in the Dry Way.—A. Carnot.—The author considers that this procedure, first recommended by Ebelmen, may be applied not merely to the separation, but also to the determination of a great number of metals. The sulphuretted hydrogen prepared by the reaction of hydrochloric acid upon iron sulphide passes through a washing bottle half full of water, is dried over calcium chloride, and acts upon the substance placed in a thin porcelain crucible heated to the required degree over a gas-burner. By heating very gently at first we may expel, without any loss of metal, in presence of the sulphurising gas the last traces of ammoniacal salts left by an imperfect washing of precipitates. At a higher temperature the simultaneous action of the hydrogen and of the sulphur vapours derived from the dissociation of the sulphuretted hydrogen perfectly sulphurises metallic oxides, carbonates, sulphates, and arseniates. The gas may thus be substituted for the use of alkaline sulphides at a high temperature.

Transformation of Hydrocellulose into Pulverulent Pyroxyles.—Aimé Girard.—Hydrocellulose placed in the same conditions as cellulose is nitrified to the same degree, yielding pyroxyle, whose composition closely agrees with hexa-nitro-cellulose.

Action of Boron Fluoride upon Aceton.—M. F. Landolph.—Aceton absorbs an equivalent of boron fluoride. The liquid thus obtained may be separated by fractional distillation into aceton fluoborides, α and β , and boraceton.

Determination of Urea.—C. Méhu.—In order to obtain in the cold all the nitrogen of the urea present in a urine, sugar must first be added to a known volume of the urine before the reaction of sodium hypobromite.

Iron Reduced by Hydrogen.—H. Moissan.—If the current of hydrogen is not dry, rapid, and long continued, and if the temperature is not uniform, the result is a mixture of iron, of ferrous and ferroso-ferric oxides.

Textile Colorist. Vol. i., No. 5, May, 1879.

This issue contains nothing of general interest.

No. 6, June, 1879.

Mr. J. H. Stebbins, of New York, has invented a new coal-tar dye, "toluol orange," the name of which indicates its source. He is also patenting a series of new dyes under the names of cresolidine, pyrogallidine, naphthylamidine, salicylidine, and picridine.

No. 7, July, 1879.

Detection of Cochineal upon the Fibre.—W. Stein recommends to boil the dyed goods under examination

with solution of sulphate of alumina and let stand for about ten minutes. If cochineal be present the liquid will turn red, but without fluorescence. If the solution is then mixed with an equal volume of solution of bisulphate of soda no discolouration should take place. Alcohol boiled with a swatch of the goods will not take up any colour.

Gazzetta Chimica Italiana.

Anno ix., 1879. Fasc. 4 and 5.

On Nicotin.—Gustavo Andreoni.—The author questions the commonly received formula.

The rest of this issue consists of extracts from other chemical journals.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 66, June, 1879.

Report presented by M. le Comte du Moncel, on behalf of the Committee of Economical Arts on M. E. Reynier's "Incandescence" Electric Lamp.—The author undertakes to combine the effects of incandescence with those resulting from the voltaic arc. He has been able to light five lamps with the current of a Bunsen battery of 30 elements, and even to keep burning for more than a quarter of an hour one of these lamps with the current of a Planté's polarisation battery of three elements.

Report by M. Debray on the Use of White Metal and Iridescent Laminæ in Embroidery.—The author uses wires of German silver in the manufacture of metallic laces. He likewise colours metallic foils for the same use by depositing upon them iridescent films of oxide of lead. He dissolves litharge in caustic alkali, and decomposes this liquid by a current of constant intensity, the positive pole of the battery being in communication with the metallic surface to be coated.

Report by M. Schützenberger on the Manufacture of Methylanilin on Vincent's Method.—This process has been already noticed.

Chemiker Zeitung.

No. 31, July 31, 1879.

The Gravimetric Determination of Fatty Matter in Milk.—Dr. F. Soxhlet.—The author proposes a special apparatus for the extraction of the dried milk-residuum with ether.

No. 32, August 7, 1879.

The Use of Ammonia Soda in the Ultramarine Manufacture.—C. Furstenau.—The author recommends the manufacturers of this soda to calcine it thoroughly, so that it may not contract further on ignition. The unequal and occasionally great shrinkage which these sodas undergo at a red heat is the cause of the fluctuating and often unsatisfactory results which they yield in the ultramarine manufacture.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 14, July 31, 1879.

Endemic Diarrhoea of Cochin-China.—According to M. Colin, santonin is a complete cure for this obstinate disease, destroying the *Anguillula* by which it is occasioned.

New Remedy for the Phylloxera.—It is said that the ravages of this insect may be checked by planting the Eucalyptus in the neighbourhood of the vineyards. An anonymous article contends from the analogy of other imported insects, such as the house-bug, the cock-roach

the American apple blight, &c., that there are no grounds for expecting the spontaneous disappearance of the phylloxera.

No. 15, August 7, 1879.

Source of Muscular Power.—In reviewing Prof. Frankland's recent memoir on this subject M. Moigno remarks that the author is wrong in not letting electricity intervene, without which the transformation of chemical action into dynamical force is impossible.

Reimann's Färber Zeitung,
No. 29, 1879.

A New Blue Dye.—Reichenbach's wood-tar colour Pittacal has been resuscitated by A. Grätzel, and it is now an article of commerce at the price of £4 per kilo, under the formidable name of "German-Imperial-Flower-Blue," with reference probably to the blue cornflower, which is said to be the favourite cognisance of the German Emperor. The pure base is insoluble in water, but dissolves in every acid, and the solutions can be diluted to any extent. The acetate is generally used for dyeing, dissolved in a little acetic acid diluted with water, and almost neutralised with ammonia. In this bath, silk and wool take a fine reddish blue without the aid of any mordant. Cotton and other vegetable fibres are prepared with a solution of tannin, followed by a solution of tartar emetic. The colours produced are perfectly fast.

Atti della R. Accademia dei Lincei.
Fascicolo I.

Distribution of Cerium, Lanthanum, and Didymium.—S. Cossa.—This memoir refers, in the first place, to apatites, which present on examination with the spectroscopic the absorption-line characteristic of didymium. The presence of cerium, lanthanum, and didymium has also been recognised in many apatites which do not present this characteristic of the absorption-spectrum. Small quantities of the same metals have also been found in limestones, in bones, and in the ashes of plants, and in the Scheelite of Traversella. The author, lastly, describes the absorption-spectra of certain didymiferous minerals.

Tungstate of Didymium.—S. Sella.—The crystals form small and imperfect octahedra of a yellow colour, shading into rose.

Preparation of Ammonialdehyds with Mixed Radicles.—R. Schiff.—The author describes the action of benzoic aldehyd upon chloral-ammonium and upon butyl-chloral-ammonium.

Chemical Nature of the Essential Oils of Laurel and of Bitter Almonds.—Prof. Fileti.—The author shows that these essences contain the nitrile—



Derivative of Chloral-ammonium.—G. Tassinari.—The compound appears in white scales, which melt at 96° to 97°, and which turn yellow and are decomposed on exposure to light. Acetyl chloride and acetic anhydride do not form acetyl derivatives, but effect a profound decomposition.

Propyl-benzoic Acid.—E. Paterno.

Certain Derivatives of Camphotymol.—E. Paterno and F. Canzoneri.

Constitution of the Cymen of Cuminic Alcohol and its Thymols.—E. Paterno and P. Spica.

Cumophenol Carbonic Acid.—E. Paterno and G. Mazzara.—These four memoirs have been already noticed.

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III. DEPARTMENT OF EVENING CLASSES.

Classes conducted by the Professors and Lecturers of the College and external Lecturers are held during the Winter Months in nearly all the Arts and Science subjects.

The NEXT SESSION will COMMENCE:—In the Arts, &c., and Medical Departments, on the 7th October; and in the Evening Classes on the 13th October. Candidates for admission must not be under 14 years of age; those under 16 will be required to pass a preliminary examination in English, Arithmetic, and Elementary Latin. Prospectuses of the several Departments may be obtained from Mr. Cornish, Piccadilly, and other Booksellers in Manchester, and at the College.

J. HOLME NICHOLSON, Registrar.

THE CHEMICAL NEWS.

VOL. XL. No. 103.

ON RADIANT MATTER.*

By WILLIAM CROOKES, F.R.S.

To throw light on the title of this lecture I must go back more than sixty years—to 1816. Faraday, then a mere student and ardent experimentalist, was 24 years old, and at this early period of his career he delivered a series of lectures on the General Properties of Matter, and one of them bore the remarkable title, *On Radiant Matter*. The great philosopher's notes of this lecture are to be found in Dr. Bence Jones's "Life and Letters of Faraday," and I will here quote a passage in which he first employs the expression *Radiant Matter* :—

"If we conceive a change as far beyond vaporisation as that is above fluidity, and then take into account also the proportional increased extent of alteration as the changes rise, we shall perhaps, if we can form any conception at all, not fall far short of Radiant Matter; and as in the last conversion many qualities were lost, so here also many more would disappear."

Faraday was evidently engrossed with this far-reaching speculation, for three years later—in 1819—we find him bringing fresh evidence and argument to strengthen his startling hypothesis. His notes are now more extended, and they show that in the intervening three years he had thought much and deeply on this higher form of matter. He first points out that matter may be classed into four states—solid, liquid, gaseous, and radiant—these modifications depending upon differences in their several essential properties. He admits that the existence of Radiant Matter is as yet unproved, and then proceeds, in a series of ingenious analogical arguments, to show the probability of its existence.†

If, in the beginning of this century, we had asked, What is a Gas? the answer then would have been that it is matter, expanded and rarefied to such an extent as to be impalpable, save when set in violent motion; invisible, incapable of assuming or of being reduced into any definite form like solids, or of forming drops like liquids; always ready to expand where no resistance is offered, and to contract on being subjected to pressure. Sixty years ago such were the chief attributes assigned to gases. Modern

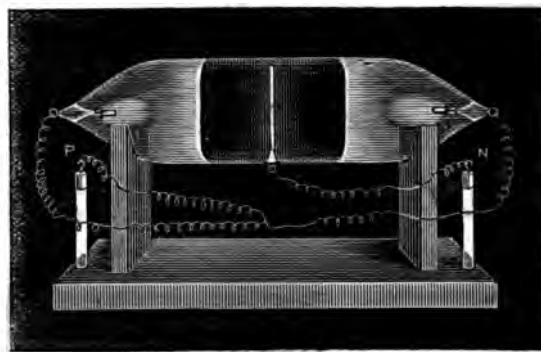
research, however, has greatly enlarged and modified our views on the constitution of these elastic fluids. Gases are now considered to be composed of an almost infinite number of small particles or molecules, which are constantly moving in every direction with velocities of all conceivable magnitudes. As these molecules are exceedingly numerous, it follows that no molecule can move far in any direction without coming in contact with some other molecule. But if we exhaust the air or gas contained in a closed vessel, the number of molecules becomes diminished, and the distance through which any one of them can move without coming in contact with another is increased, the length of the mean free path being inversely proportional to the number of molecules present. The further this process is carried the longer becomes the average distance a molecule can travel before entering into collision; or, in other words, the longer its mean free path, the more the physical properties of the gas or air are modified. Thus, at a certain point, the phenomena of the radiometer become possible, and on pushing the rarefaction still further, *i.e.*, decreasing the number of molecules in a given space and lengthening their mean free path, the experimental results are obtainable to which I am now about to call your attention. So distinct are these phenomena from anything which occurs in air or gas at the ordinary tension, that we are led to assume that we are here brought face to face with Matter in a Fourth state or condition, a condition as far removed from the state of gas as a gas is from a liquid.

Mean Free Path. Radiant Matter.

I have long believed that a well-known appearance observed in vacuum tubes is closely related to the phenomena of the mean free path of the molecules. When the negative pole is examined while the discharge from an induction-coil is passing through an exhausted tube, a dark space is seen to surround it. This dark space is found to increase and diminish as the vacuum is varied, in the same way that the mean free path of the molecules lengthens and contracts. As the one is perceived by the mind's eye to get greater, so the other is seen by the bodily eye to increase in size; and if the vacuum is insufficient to permit much play of the molecules before they enter into collision, the passage of electricity shows that the "dark space" has shrunk to small dimensions. We naturally infer that the dark space is the mean free path of the molecules of the residual gas, an inference confirmed by experiment.

I will endeavour to render this "dark space" visible to all present. Here is a tube (Fig 1), having a pole in the

FIG. 1.



centre in the form of a metal disk, and other poles at each end. The centre pole is made negative, and the two end poles connected together are made the positive terminal. The dark space will be in the centre. When the exhaustion is not very great the dark space extends only a little on each side of the negative pole in the centre. When the exhaustion is good, as in the tube before you, and I

* A Lecture delivered to the British Association for the Advancement of Science, at Sheffield, Friday, August 22, 1879.

† "I may now notice a curious progression in physical properties accompanying changes of form, and which is perhaps sufficient to induce, in the inventive and sanguine philosopher, a considerable degree of belief in the association of the radiant form with the others in the set of changes I have mentioned."

"As we ascend from the solid to the fluid and gaseous states, physical properties diminish in number and variety, each state losing some of those which belonged to the preceding state. When solids are converted into fluids, all the varieties of hardness and softness are necessarily lost. Crystalline and other shapes are destroyed. Opacity and colour frequently give way to a colourless transparency, and a general mobility of particles is conferred."

"Passing onward to the gaseous state, still more of the evident characters of bodies are annihilated. The immense differences in their weight almost disappear; the remains of difference in colour that were left, are lost. Transparency becomes universal, and they are all elastic. They now form but one set of substances, and the varieties of density, hardness, opacity, colour, elasticity and form, which render the number of solids and fluids almost infinite, are now supplied by a few slight variations in weight, and some unimportant shades of colour."

"To those, therefore, who admit the radiant form of matter, no difficulty exists in the simplicity of the properties it possesses, but rather an argument in their favour. These persons show you a gradual resignation of properties in the matter we can appreciate as the matter ascends in the scale of forms, and they would be surprised if that effect were to cease at the gaseous state. They point out the greater exertions which Nature makes at each step of the change, and think that, consistently, it ought to be greatest in the passage from the gaseous to the radiant form."—*Life and Letters of Faraday*, vol. i., p. 308.

turn on the coil, the dark space is seen to extend for about an inch on each side of the pole.

Here, then, we see the induction spark actually illuminating the lines of molecular pressure caused by the excitement of the negative pole. The thickness of this dark space is the measure of the mean free path between successive collisions of the molecules of the residual gas. The extra velocity with which the negatively electrified molecules rebound from the excited pole keeps back the more slowly moving molecules which are advancing towards that pole. A conflict occurs at the boundary of the dark space, where the luminous margin bears witness to the energy of the discharge.

Therefore the residual gas—or, as I prefer to call it, the gaseous residue—within the dark space is in an entirely different state to that of the residual gas in vessels at a lower degree of exhaustion. To quote the words of our last year's President, in his Address at Dublin:—

"In the exhausted column we have a vehicle for electricity not constant like an ordinary conductor, but itself modified by the passage of the discharge, and perhaps subject to laws differing materially from those which it obeys at atmospheric pressure."

In the vessels with the lower degree of exhaustion, the length of the mean free path of the molecules is exceedingly small as compared with the dimensions of the bulb, and the properties belonging to the ordinary gaseous state of matter, depending upon constant collisions, can be observed. But in the phenomena now about to be examined, so high is the exhaustion carried that the dark space around the negative pole has widened out till it entirely fills the tube. By great rarefaction the mean free path has become so long that the hits in a given time in comparison to the misses may be disregarded, and the average molecule is now allowed to obey its own motions or laws without interference. The mean free path, in fact, is comparable to the dimensions of the vessel, and we have no longer to deal with a *continuous* portion of matter, as would be the case were the tubes less highly exhausted, but we must here contemplate the molecules *individually*. In these highly exhausted vessels the molecules of the gaseous residue are able to dart across the tube with comparatively few collisions, and radiating from the pole with enormous velocity, they assume properties so novel and so characteristic as to entirely justify the application of the term borrowed from Faraday, that of *Radiant Matter*.

Radiant Matter exerts Powerful Phosphorogenic Action where it Strikes.

I have mentioned that the Radiant Matter within the dark space excites luminosity where its velocity is arrested by residual gas outside the dark space. But if no residual gas is left, the molecules will have their velocity arrested by the sides of the glass; and here we come to the first and one of the most noteworthy properties of Radiant Matter discharged from the negative pole—its power of exciting phosphorescence when it strikes against solid matter. The number of bodies which respond luminously to this molecular bombardment is very great, and the resulting colours are of every variety. Glass, for instance, is highly phosphorescent when exposed to a stream of Radiant Matter. Here (Fig. 2) are three bulbs composed

of different glass: one is uranium glass (a), which phosphoresces of a dark green colour; another is English glass (b), which phosphoresces of a blue colour; and the third (c) is soft German glass,—of which most of the apparatus before you is made,—which phosphoresces of a bright apple-green.

My earlier experiments were almost entirely carried on by the aid of the phosphorescence which glass takes up when it is under the influence of the radiant discharge; but many other substances possess this phosphorescent power in a still higher degree than glass. For instance, here is some of the luminous sulphide of calcium prepared according to M. Ed. Becquerel's description. When the sulphide is exposed to light—even candlelight—it phosphoresces for hours with a bluish white colour. It is, however, much more strongly phosphorescent to the molecular discharge in a good vacuum, as you will see when I pass the discharge through this tube.

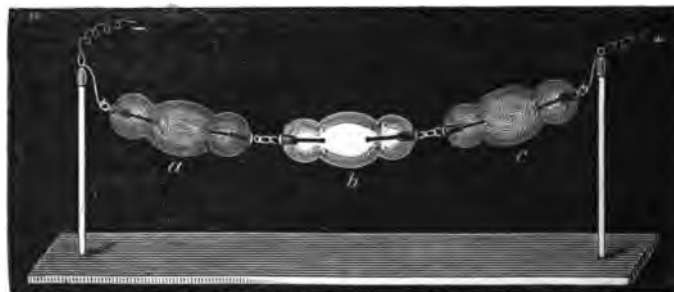
Other substances besides English, German, and uranium glass, and Becquerel's luminous sulphides, are also phosphorescent. The rare mineral Phenakite (aluminate of glucinum) phosphoresces blue; the mineral Spodumene (a silicate of aluminium and lithium) phosphoresces a rich golden yellow; the emerald gives out a crimson light. But without exception, the diamond is the most sensitive substance I have yet met for ready and brilliant phosphorescence. Here is a very curious fluorescent diamond, green by daylight, colourless by candlelight. It is mounted in the centre of an exhausted bulb (Fig. 3), and

FIG. 3.



the molecular discharge will be directed on it from below upwards. On darkening the room you see the diamond

FIG. 2.



as much light as a candle, phosphorescing of ten. the diamond the ruby is one of the most re- tones for phosphorescing. In this tube (Fig. 4)

FIG. 4.



lection of ruby pebbles. As soon as the ink is turned on you will see these rubies th a brilliant rich red tone, as if they were t. It scarcely matters what colour the ruby is, th. In this tube of natural rubies there are ll colours—the deep red and also the pale pink re are some so pale as to be almost colourless, f the highly-prized tint of pigeon's blood; but mpa&t of Radiant Matter they all phosphoresce the same colour.

ruby is nothing but crystallised alumina with puring-matter. In a paper by Ed. Becquerel,* twenty years ago, he describes the appearance as glowing with a rich red colour in the phos- Here is some precipitated alumina prepared careful manner. It has been heated to white- you see it also glows under the molecular with the same rich red colour.

trum of the red light emitted by these varieties is the same as described by Becquerel twenty There is one intense red line, a little below B in the spectrum, having a wave-length of . There is a continuous spectrum beginning at and a few fainter lines beyond it, but they are comparison with this red line that they may be This line is easily seen by examining with a at spectroscopie the light reflected from a good

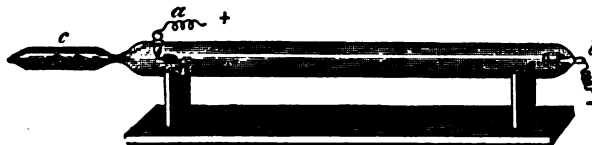
one particular degree of exhaustion more than any other for the development of the pro- Radiant Matter which are now under examina- ghtly speaking it may be put at the millionth of here.† At this degree of exhaustion the phos- is very strong, and after that it begins to ntil the spark refuses to pass.‡

de Chimie et de Physique, 3rd series, vol. lviii., p. 50, 1859
10 millionth of an atmosphere = 0.000076 millim.
5789 millionths of an atmosphere = 1.0 millim.
" " " = 760.0 millims.
" " " = 1 atmosphere.

100 years ago Mr. Wm. Morgan communicated to the ty a Paper entitled "Electrical Experiments made to Non-conducting Power of a Perfect Vacuum, &c." The trades from this Paper, which was published in the *Phil.* 785 (vol. lxxv., p. 272), will be read with interest:— rial gage about 15 inches long, carefully and accurately ery particle of air was expelled from the inside, was coated 3 inches down from its sealed end, and being inverted through a perforation in the brass cap which covered the cistern; the whole was cemented together, and the used from the inside of the cistern through a valve in , which producing a perfect vacuum in the gage formed at peculiarly well adapted for experiments of this kind. g thus adjusted (a small wire having been previously fixed le of the cistern to form a communication between the ad the mercury, into which the gage was inverted) the ras applied to the conductor of an electrical machine, and

I have here a tube (Fig. 5) which will serve to illustrate the dependence of the phosphorescence of the glass on the

FIG. 5.



degree of exhaustion. The two poles are at *a* and *b*, and at the end (*c*) is a small supplementary tube connected with the other by a narrow aperture, and containing solid caustic potash. The tube has been exhausted to a very high point, and the potash heated so as to drive off moisture and injure the vacuum. Exhaustion has then been re-commenced, and the alternate heating and exhaustion repeated until the tube has been brought to the state in which it now appears before you. When the induction spark is first turned on nothing is visible—the vacuum is so high that the tube is non-conducting. I now warm the potash slightly and liberate a trace of aqueous vapour. Instantly conduction commences, and the green phosphorescence flashes out along the length of the tube. I continue the heat, so as to drive off more gas from the potash. The green gets fainter, and now a wave of cloudy luminosity sweeps over the tube, and stratifications appear, which rapidly get narrower, until the spark passes along the tube in the form of a narrow purple line. I take the lamp away, and allow the potash to cool; as it cools, the aqueous vapour, which the heat had driven off, is re-absorbed. The purple line broadens out, and breaks up into fine stratifications; these get wider, and travel towards the potash tube. Now a wave of green light appears on the glass at the other end, sweeping on and driving the last pale stratification into the potash; and now the tube glows over its whole length with the green phosphorescence. I might keep it before you, and show the green growing fainter and the vacuum becoming non-conducting; but I should detain you too long, as time is required for the absorption of the last traces of vapour by the potash, and I must pass on to the next subject.

(To be continued.)

Use of the Diffusion Method in the Study of the Phenomena of Dissociation.—L. Troost.—It has been already established that diffusion cannot solve the question of the existence or non-existence of bodies in the state of definite gaseous compounds. The experiments of M. A. Naumann and of MM. Wiedemann and Schulze can in no manner solve the question of the existence or non-existence of chloral hydrate as a definite gaseous compound. —*Comptes Rendus*.

notwithstanding every effort, neither the smallest ray of light, nor the slightest charge, could ever be procured in this exhausted gage."

"If the mercury in the gage be imperfectly boiled, the experiment will not succeed; but the colour of the electric light, which in air rarefied by an exhauster is always violet or purple, appears in this case of a beautiful green, and, what is very curious, the degree of the air's rarefaction may be nearly determined by this means; for I have known instances, during the course of these experiments, where a small particle of air having found its way into the tube, the electric light became visible, and as usual of a green colour; but the charge being often repeated, the gage has at length cracked at its sealed end and in consequence the external air, by being admitted into the inside, has gradually produced a change in the electric light from green to blue, from blue to indigo, and so on to violet and purple, till the medium has at length become so dense as no longer to be a conductor of electricity. I think there can be little doubt, from the above experiments, of the non-conducting power of a perfect vacuum."

"This seems to prove that there is a limit even in the rarefaction of air, which sets bounds to its conducting power; or, in other words, that the particles of air may be so far separated from each other as no longer to be able to transmit the electric fluid; that if they are brought within a certain distance of each other, their conducting power begins, and continually increases till their approach also arrives at its limit."

REPORT OF THE
COMMITTEE ON THE CHEMISTRY OF
SOME OF THE LESSER-KNOWN ALKALOIDS,
ESPECIALLY VERATRIA AND BEBEERINE.*

SINCE last year investigations have been made on the alkaloids contained in *Veratrum album*, and *V. viride*, with the following general results; as the details of the experiments have already been communicated to the Chemical Society in two papers (*Journal of the Chemical Society*, 1879, i., pp. 405 and 421), it is unnecessary to quote them here.

Each kind of root was treated by the process described in last year's Report, viz., percolating with alcohol acidulated with tartaric acid, evaporating to a small bulk, treating with water to precipitate resin, filtering, alkalinising with soda, and repeatedly shaking with a large bulk of ether, the ethereal solutions of alkaloids, &c., thus obtained being agitated with aqueous tartaric acid to remove the bases and then used over again. In each case a certain amount of flocculent alkaloidal matter was left undissolved by the ether, consisting mainly of an alkaloid analogous to jervine, but differing therefrom in certain respects, to which accordingly the name *Pseudojervine* is applied. The solutions of tartrates of alkaloids obtained were treated with soda and about an equal bulk of ether, whereby a large portion of the bases was dissolved in each case, but some left undissolved, especially with the *V. album* product; this insoluble matter contained pseudojervine, together with a little jervine, and in the case of the *V. album* product, a large quantity of an uncrystallisable base sparingly soluble in ether, to which the term *Veratralbine* is applied, as this body does not seem to be present in *V. viride* roots in any considerable proportion. The second ethereal solutions thus obtained deposited in each case crystals of jervine and a little of a new base to which the term *Rubijervine* is applied; the mother-liquors of these crystals dried up to varnish-like masses, which were not identical in the two cases; the product from *V. album* roots consisted essentially of veratralbine, with a minute quantity of an alkaloid forming veratric acid on saponification with alcoholic potash; this base was the only alkaloid of the saponifiable class present in the roots; presumably it was the *veratrine* obtainable from *V. sabadilla* seeds, as described in last year's Report, inasmuch as the mixture of this base and veratralbine obtained was powerfully sternutatory, whilst the peculiar tendency to provoke sneezing was lost on treatment with alcoholic potash (neither jervine, pseudojervine, rubijervine, nor veratralbine produces sneezing). The product from the *V. viride* roots was even more powerfully sternutatory than that from the *V. album* roots; it consisted, however, almost wholly of *Cevadine* (the second crystallisable alkaloid obtainable from *V. sabadilla* seeds, as described in last year's Report), not more than traces of either veratralbine or veratrine being contained; on saponification it yielded about the theoretical quantity of cevadic acid (the *methyl-crotonic acid* of Frankland and Duppa, identical with the *tiglic acid* of Geuther).

The following table shows the approximate quantities of the different alkaloids contained in a kilogramme of each root examined:—

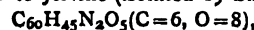
	<i>V. album</i> .	<i>V. viride</i> .
Jervine.. ..	1·3 grammes	0·2 gramme
Pseudojervine	0·4 "	0·15 "
Rubijervine..	0·25 "	0·02 "
Veratralbine	2·2 "	Not more than traces
Veratrine ..	0·05 "	Trace; less than 0·004
Cevadine ..	Apparently absent	0·43 gramme
	4·20	0·80

* Read before Section B, British Association for the Advancement of Science, Sheffield Meeting, August 21, 1879. The Committee consisted of Mr. W. Chandler Roberts, F.R.S. (Secretary), Dr. C. R. Alier Wright, and Mr. A. P. Luff.

The *V. album* roots were consequently about five times as rich in total alkaloids as the *V. viride* roots.

The following are the chief characteristics and properties of the new alkaloids examined; in many respects the statements of former observers concerning the alkaloids of these two kinds of roots appear to be erroneous, probably owing to the complete separation of jervine, &c., from the other substances now shown to be also present never having been previously effected.

Jervine.—When crystallised, $C_{26}H_{37}NO_3 \cdot 2H_2O$: if the crystals separate from too hot or concentrated alcoholic liquors, somewhat less water is frequently present; readily becomes anhydrous at 100°; melts at 237° to 239° (purest specimens—corrected). Forms an almost insoluble sulphate, and a very sparingly soluble hydrochloride and nitrate. The gold salt is $C_{26}H_{37}NO_3 \cdot HCl \cdot AdCl_3 \cdot H_2O$, the water of crystallisation being lost only slowly at 100°. With strong sulphuric acid dissolves to a yellow fluid, quickly darkening to a greenish brown, which soon becomes a fine green by absorption of a little water from the air if in an open dish; if in a test-tube, becomes green by cautiously adding minute quantities of water. Not sternutatory: not saponifiable. The formula assigned in 1837 by Will to jervine (isolated by Simon),—



modified by Gerhardt and his followers to $C_{30}H_{46}N_2O_5$, is considerably incorrect, the error being apparently due to an imperfect nitrogen determination (by volume), and to the presence of pseudojervine in the substance examined.

Pseudojervine.—Crystallises anhydrous, $C_{25}H_{43}NO_7$. Externally resembles jervine closely: melts at 299° (corrected: forms a sulphate crystallisable and soluble in water, especially when hot. Hydrochloride very sparingly soluble in water even when hot, provided no free hydrochloric acid is present. Gives with sulphuric acid exactly the same colour reaction as jervine. Not saponifiable: not sternutatory.

Rubijervine.—Crystallises anhydrous, $C_{26}H_{43}NO_2$: resembles jervine in appearance, and melts at nearly the same temperature (236° purest specimen—corrected). Sulphate and hydrochloride crystallisable and readily soluble in water, especially if warm. With strong sulphuric acid forms a yellow solution, becoming brownish yellow, brownish orange, brownish blood-red, and ultimately brownish purple by absorption of moisture: by cautious dilution with water the brownish blood-red fluid becomes successively crimson, purple, dark lavender, dark violet, light indigo. Not saponifiable: not sternutatory.

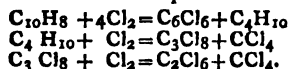
Veratralbine.—Amorphous, approximately $C_{26}H_{43}NO_5$. No crystallisable salts obtained as yet. With sulphuric acid dissolves to a yellow fluid, becoming brownish orange and brownish blood-red, with a strong green fluorescence; in this respect it closely resembles cevadine, which only differs in giving somewhat clearer tints, a crimson-magenta coloured fluid of a peculiarly beautiful and permanent shade being developed on absorption of a trace of moisture; veratrine (of Couerbe) gives precisely the same colours as cevadine, but the dark red solution formed before the crimson tint is developed by absorption of moisture does not exhibit any fluorescence. Veratralbine is not saponifiable, and is not sternutatory.

Currents of Ampère.—M. Trève.—Ampère asked if the molecular currents of magnets are entirely created in the magnetic substance during magnetisation, or if the magnetising cause merely determines a circulation of currents pre-existing in the metals in their natural state. He inclined to the latter opinion. The author thinks that there is every reason to admit, with Ampère, that the particular currents pre-exist in the magnetic metals, and that the current of the battery merely determines the circulation and the direction.—*Comptes Rendus*.

RESULTS OF AN EXHAUSTIVE CHLORINATION
OF ISO-DINAPHTHYL.

By WATSON SMITH, F.C.S., F.I.C.

THE investigations of Ruoff and Merz have already shown that an exhaustive chlorination of naphthalene by continuous strong heating in sealed tubes with antimony pentachloride, or of perchlor-naphthalene with either antimony pentachloride or chloride of iodine, gives rise to a disintegration of the naphthalene molecule, so that perchlor-benzene, with perchlor-ethane and perchlor-methane, are formed. The formation of the two latter perchlorinated products is explained by assuming with Graebe (*Ann. Chem. Pharm.*, 149, 21) that the group $(C_4H_4)''$ contained in the naphthalene molecule, together with the benzene nucleus, is converted into perchlor-butane, which immediately after its formation, and at the high temperature, splits up into perchlor-propane and perchlor-methane, and the former again into perchlor-ethane and perchlor-methane. Thus:—



(See Kraft and Merz, *Berichte der Deutsch.*, 1875, 1300.)

Diphenyl by exhaustive chlorination yields perchlor-diphenyl—



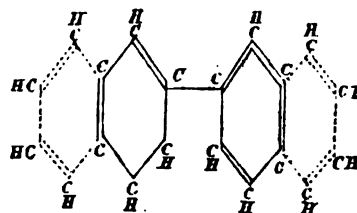
very difficult to sublime, giving feathery needles, which did not melt even at 270° . The chlorination of a large number of the most prominent members of the aromatic series by Merz in conjunction with others (Ruoff, Moe, &c.) has done perhaps more than any other line of research to demonstrate the truth involved in Kekulé's theory, by which all aromatic compounds are represented as containing the benzene nucleus, for by this method of exhaustive chlorination from every aromatic body treated (excepting diphenyl) perchlor-benzene has been obtained, and with regard to the by-perchlorinated products (perchlor-paraffin groups), these have also indicated the truth of the mode of the constitution-formulae adopted.

It was now thought interesting to attempt the perchlorination or exhaustive chlorination of iso-dinaphthyl. For this purpose the hydrocarbon, after being introduced into the perfectly dry tube, was treated with ten times its weight of antimony pentachloride, gradually introduced down a long funnel. The whole mass of the dinaphthyl was immediately turned black and frothed up strongly, evolving volumes of hydrochloric acid gas. When the whole of the pentachloride had been added, the tube was sealed up and heated first to a temperature of 150° for some two to three hours. On cooling and opening, a considerable pressure was manifested, resulting from accumulation of hydrochloric acid.

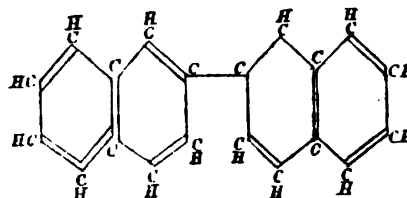
A stream of dry chlorine was now passed through the mass to convert all the trichloride formed into pentachloride. The tube was then again sealed up, and heated some two to three hours to 250° . On opening, still considerable pressure, but less than before. Again chlorinated, sealed, and heated to 350° . But little pressure, and on again heating to some 400° , after re-chlorinating, and re-opening, no pressure perceptible. The product—after well washing with excess of concentrated hydrochloric acid, and then with warm tartaric acid solution, and finally with water, till no more trace of acidity remained, by which all antimony compound was removed—was observed to be rather resinous. On sublimation of a portion a quantity of small needles was obtained, having an odour somewhat like camphor, and melting at about 182° , evidently perchlorethane. Perchlormethane was also most probably present in the oily portion. The mass was now once more treated with ten times its weight of antimony pentachloride. Further action immediately set in, with evolution of hydrochloric acid, proving that full perchlorination

was prevented before by the protecting coating of antimony chlorides formed. After sealing up, and heating exactly as described above, a mass of long needles was observed as the final product. On washing as before, drying at 80° , and subliming, first small needles or prisms of prisms of perchlorethane, with odour of camphor, and melting-point of 182° , were obtained, succeeded by a large quantity of fine long needles melting at 220° to 223° , and evidently consisting of perchlor-benzene (C_6Cl_6). It is therefore proved that isodinaphthyl does not yield perchlor-diphenyl, or rather, that the diphenyl residue, which may be supposed to exist in dinaphthyl, suffers disruption in the process of perchlorination.

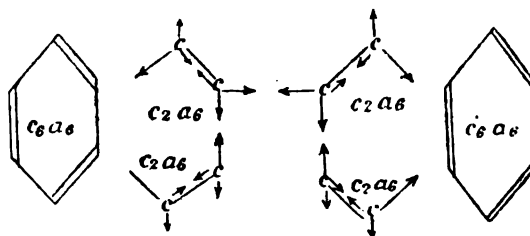
This is an interesting point, because Ruoff and Merz found that the diphenyl group alone, i.e., diphenyl itself, could not be broken up by perchlorination, but yielded simply $C_{12}Cl_{10}$, perchlor-diphenyl. This diphenyl residue in iso-dinaphthyl can be rendered visible in the following formula:—



As I was not able to detect perchlor-methane in the products obtained, though I by no means assert it was not formed in the reaction to some extent, I must for the present consider that in all probability iso-dinaphthyl breaks up evenly into perchlor-benzene and perchlor-ethane, thus:—



yields



Or $C_{10}H_7 - C_{10}H_7$ yields $2C_6Cl_6 + 4C_2Cl_6$.

It would next be interesting to try the effect of exhaustive chlorination on the other two dinaphthyl isomerides in order to see if the different mode of binding together will denote a different mode of disruption of the molecule. The exhaustive chlorination of the new hydrocarbon phenyl-naphthalene, lately synthesised by me, may very probably give perchlor-diphenyl together with perchlorethane, or a mixture of perchlor-ethane and perchlor-methane. It seems very probable that on the one side the simple benzene nucleus may maintain a kind of equilibrium with the composite benzene nucleus on the other side of the link, and thus a perchlorinated diphenyl residue will remain, whilst the residue, $(C_4H_4)''$, will be split

off into perchlorinated ethane, or methane and ethane groups. The solution of these latter problems will occupy me next, and I hope soon to be able to give a further report on the subject. I ought also to mention that after the first stage in the isodinaphthyl chlorination, when I obtained a resinous product, I dried a portion of this, after well washing as detailed, and on sublimation obtained crystals of perchlorethane, but not at all abundantly. A portion of the body dissolved in petroleum spirit, boiled with animal charcoal, filtered, and evaporated to dryness, gave apparently small round warty masses, which the magnifying glass showed to consist of stellate prismatic groups. It is possible this body may be perchlor-isodinaphthyl, and this possibility I shall investigate, as also that of the formation of perchlor-phenyl-naphthalene.

University Laboratory, Zürich,
August 11, 1879.

IS OZONE PRODUCED DURING THE ATMOSPHERIC OXIDATION OF PHOSPHORUS?

By C. T. KINGZETT.

I AM chiefly induced to make the present brief communication from a conviction that most chemists believe and teach at the present time that ozone is produced during the atmospheric oxidation of phosphorus, just as a time ago it was believed and taught that the same substance is produced during the atmospheric oxidation of turpentine and other terpenes. I should, however, have deferred any communication of my own upon the subject until I might have had a better opportunity of practically investigating it, but for reading several interesting papers by Mr. A. R. Leeds, which have been printed in these pages from time to time.

On the face of things, it appears to me that one would not expect ozone to be formed by the aerial oxidation of phosphorus now that we understand the constitution of ozone, and since peroxide of hydrogen is the only known agent which resembles ozone in its general properties, and since, further, peroxide of hydrogen is now known to be produced in various processes of slow oxidation, it is this substance which should be more naturally expected to be produced in connection with the oxidation of phosphorus.

In a paper on a "New Reaction for Iodates and Iodides," M. Corne states* that the water in which phosphorus has been kept for some time liberates iodine from solutions of these substances, and he, as I think, illogically concludes that this behaviour is due to the presence of phosphorous acid. As a matter of fact, the reaction is characteristic of peroxide of hydrogen. This is important, if it be remembered that ozone is not soluble to any considerable extent in water, and that this is the only other substance which, as a product of the oxidation of phosphorus, could give the same reaction. This test, however, is not conclusive as regards the nature of the active agent, since there is a controversy as to the solubility of ozone in water. It will be remembered Marignac demonstrated that the formation of the active agent requires the presence of water, and this in itself is a further reason for regarding it as peroxide of hydrogen.

Mr. A. R. Leeds,† who regards the active agent as ozone and writes of it as such, has described an apparatus by the use of which it may be more freely obtained than usual, and he finds that not only is it produced by the passage of air or oxygen over phosphorus partially submerged in water, but that it may also be obtained, and in greater quantity too, by using an acidulated solution of potassic dichromate instead of water. He found that the best solution to employ was one containing in every 1250 c.c. 150 c.c. H_2SO_4 and 25 grms. of the dichromate. By con-

necting the vessels containing the phosphorus and this that the amount of active agent generated in the first of a mixture by appropriate tubes, Mr. Leeds ascertained, series of such vessels is increased by causing the air containing it to pass through the second and third vessels and so on. The temperature necessary for maximum production he fixed at about 24°C . Operating in this way, he obtained a gaseous product containing about 2.5 milligrams of estimated ozone (?) to the litre of air.

The estimations were made by first washing the air in water, and then passing it into a solution of potassic iodide, determining the iodine thus set free by means of a standard solution of sodic hyposulphite. Now, although at first blush it might be thought that the washing water would remove any peroxide of hydrogen which might be held in the gas in the vesicular state, yet as a matter of fact this is by no means necessarily the case. Indeed, all the erroneous views which were once prevalent upon the nature of antozone were due to the same misleading notion. Antozone is now known to be peroxide of hydrogen, but from the fact that it was obtained diffused through large volumes of air, and that ordinary washing with water did not effectually remove it, it was regarded for a long time as a new substance. It is, therefore, I submit more than probable that the washing with water to which Mr. Leeds subjected his gaseous product did not remove the peroxide of hydrogen therefrom, and so he estimated it as, and called it, ozone.

Not rarely Andrew's experiments are referred to as establishing the idea of ozone being a product of the oxidation of phosphorus. But if his research be carefully studied, it will be observed that he gives no proof of the ozonic nature of his product. He merely showed that his active air obtained from phosphorus lost its properties when agitated with water, suggesting to my mind that these were due to peroxide of hydrogen which the water abstracted during the experiment.

Most practical chemists are aware of the great difficulty experienced in perfectly separating two gases by passing them through a solution in which one of them is soluble. For instance, it has been noticed (and more particularly, I believe, by Sir B. C. Brodie in some of his later researches) that when a mixture of carbonic oxide and carbonic anhydride gases is passed through a solution of potash in order to remove the latter one, the process is ineffectual except in those cases where some special precautions are observed. To be effectual it is requisite that the potash solution should present a very large surface to the mixed gases, such as is only obtained by conducting them through a long tube filled with the fluid. The least reflection on the nature of a gas will satisfy the mind as to the necessity of this proceeding for bringing every particle of the gaseous mixture into intimate contact with the fluid.

In addition to the objections which I have already pointed out to viewing the active agent produced in the atmospheric oxidation of turpentine as ozone, there are other considerations which lead to the other conclusion, viz., that the active agent is peroxide of hydrogen. *There is no known process of slow oxidation which has been established to produce ozone.*

In the several instances which have been written about from time to time the different observers have always relied upon properties common to the two substances, and have never instituted volumetric investigations which are alone sufficient to decide the question. On the other hand, there are known several processes of slow oxidation in which peroxide of hydrogen is formed, as, for instance, those relating to ether and the terpenes; and I believe that as peroxide of hydrogen is formed in each of these cases as a secondary product due to the action of water upon a peroxide, so also the oxidation of phosphorus by air gives rise to an oxide which generates peroxide of hydrogen by contact with water. So far as I remember

* *Journ. de Pharm.*, [4], xxii., 425.

† *CHEMICAL NEWS*, vol. xxxix., p. 157; also vol. xl., p. 70.

* In his last paper he identifies this substance among the other products.

† *Philosophical Transactions*, 1856, p. 1.

the experiments of D. J. Boche* upon this last-named subject, they bear out my view. At any rate the active agent produced in the aerial oxidation of phosphorus has never been proved to possess the special volumetric relations of ozone, and until this is done it seems to me a pity that in chemical text-books such very decided statements should be made as are to be found.

ON THE COMPOSITION OF A WELL-WATER AT GROUVILLE WHICH OCCASIONED AN OUTBREAK OF TYPHOID FEVER.

By THOS. M. MORGAN, F.I.C., &c.

THE Female Orphan's Home of Grouville, one of the parishes on the eastern part of this island, was visited in the latter end of last May by a severe outbreak of typhoid fever: out of nearly one hundred and fifty inmates no fewer than twenty-five were prostrated by the disease.

The establishment is situated in the country, pretty well isolated, and far removed from any sewers; there was, moreover, no fever in the neighbourhood, and therefore the source of this was probably to be found on the spot. The well yielded water of a pleasant taste, perfectly clear and colourless, and much esteemed; it therefore did not lie under much suspicion, but was nevertheless sent to me for analysis. It contained neither nitrites nor ammonia, and decolourised very little permanganate. Analysis showed per million parts:—

Total Solids.	Chlorine.	N as Nitrites.	C Organic I. II.	O of KMnO ₄ consumed.
1378	326.6	26	2 1.8	0.186

The dissolved gases per litre were:—

CO ₂ .	O.	N.
58 c.c.	4 c.c.	15 c.c.

The solid residue was highly deliquescent, from the abundance of calcium nitrate present; it also contained much sodium chloride, and the carbonates of lime and magnesia.

The determination of chlorine in the waters here does not afford much information in respect of sewage contamination, as nearly all the wells, even though much above the sea-level, contain it in abundance; the quantity in this is, however, nearly twice as great as in a new well made a hundred yards distant, the analysis of which is given below. The nitrogen was determined by Frankland's method, except that the chlorine was not previously abstracted: the nitric oxide obtained from two or three samples of this water, and in this way was found to be quite pure. The carbon was determined by an organic analysis of the residue and the dissolved gases by Bunsen's method.

The quantity of nitrates here found was remarkable, and so far as the analysis is concerned they alone distinctly indicated a bad water. Tested as it came from the pump with ferrous sulphate and sulphuric acid, the usual indication of their presence might be indistinctly seen. Still, many waters here contain nitrites and nitrates often in considerable quantity. When kept for a few days it acquired an odour, and when this was rendered more sensible by warming in a flask, it at once suggested to me the smell of putrid urine. Another quantity, kept in a clear glass bottle and exposed to the light, gave rise to a slight deposit of green matter, which under the microscope was seen to consist of detached green cells.

The water was condemned, and the inmates of the Home were told to abstain from drinking it. Subsequently several fresh cases of fever occurred, but the patients acknowledged that, notwithstanding the prohibition, they had still used the old well. One of the older girls, who had carefully

kept the younger ones from it, ventured to take some herself, but was seized with the disease and died of it. The pump was then removed, and after that there were no more attacked, nor were there any sufferers among twelve young children fed solely on milk. Altogether there have been thirty-nine cases.

The soil in this locality consists of a yellow loam from 15 to 20 feet thick, and below that there is about an equal thickness of coarse yellow sea-sand or gravel, by sinking wells into which water is obtained.

This particular well was sunk in the courtyard of the establishment, and afterwards covered in. About 60 feet distant a cesspool had been constructed, 6 feet deep. It had long been used, but for the last three years had received urine and soap-suds exclusively. After the outbreak of fever it was emptied and filled up: from this the well is supposed to have received its pollution. Last winter, after some heavy rains, it became very muddy, and was accordingly pumped out and cemented part of the way down; perhaps this interference facilitated the entrance of impurities. Two bore wells within 25 feet of this one yielded waters containing much ammonia and nitrites. A bore well 100 yards from the cesspool, and in another direction, gave water of the following composition per million parts:—

Total Solids.	Chlorine.	Ammonia.	N as Nitrites and Nitrites.	C Organic. I. II.
840	180	traces	5.5	2.9 2.8

Dissolved gases in a litre:—

Carbon Dioxide.	Oxygen.	Nitrogen.
32.5 c.c.	3.5	13

A later analysis, when the water had been in use for some time, and when the completion of the well had exposed it more to the atmosphere, showed the quantities of carbon and nitrites had slightly decreased, and a corresponding increase in the nitrates.

It is important to note—as Dr. Bull, the consulting physician to the establishment, remarks, and he is thoroughly conversant with its medical history from the beginning—that there has never been a decided case of typhoid there before, and hence, as the sewage cannot have proceeded from typhoid patients, it is possible for this disease to arise *de novo*.

The Laboratory, Victoria College, Jersey.

NEW PROCESS FOR THE RAPID ESTIMATION OF PURE SUGAR IN RAW AND REFINED COMMERCIAL SUGARS.*

By P. CASAMAJOR.

(Continued from page 76.)

PART II.

I HAD given up hope of success, when on passing before No. 71, Maiden Lane, I saw at a window a number of bottles, labelled *Diamond Methyl*, which proved to be methylic alcohol. I determined to try if methylic alcohol would answer better than the ethylic, and purchased a gallon of the product. This stood at 92.5° by the alcoholometer, and even at this strength the gummiest sugar could be cleansed by mixing with this spirit and stirring the mixture. On saturating it with sugar, and placing the solution in the tube of a Ventzke saccharometer, the indication was 1.7, which is less than corresponds to ethylic alcohol of the same density.

These first trials were very encouraging, and those that followed were not less so. Without recounting now the experiments that were made, I may state that, by the use of methylic alcohol, I finally succeeded in obtaining, with

* Deutsch. Chem. Ges. Ber., vi., 439.

* Read before the American Chemical Society, June 5, 1879.

an alcohometer, results that agree very closely with those of the optical saccharometer, with cane sugars of all classes, from the highest to the lowest.

The explanations already given concerning the use of ethylic alcohol of various strengths, saturated with sugar, and the effects that water and the soluble impurities have in lowering their alcohometric degrees, render superfluous any account of similar experiments with hydrate of methyl. I may briefly state that, after making a great number of trials, I found that methylic alcohol of $83\frac{1}{4}^{\circ}$ of the alcohometer,* when saturated with sugar, stands at 77.1° , and that this solution is the one that has given the most accurate results.

This saturated solution is quite easily obtained by taking methylic alcohol standing at $83\frac{1}{4}^{\circ}$ by the alcohometer, and saturating it with sugar, by the process which Numa Grar suggested to Payen. This consists in shaking up the alcohol with powdered sugar. Grar made his powdered sugar by grinding it in a mortar in presence of alcohol. I have generally used dry *extra-powdered* sugar. When this cannot be had, grinding in a mortar is an excellent way of getting the sugar in fine particles, which are very necessary for obtaining a saturated solution in a short time. When very finely powdered sugar is shaken up with either ethylic or methylic alcohol, the finest particles are dissolved almost immediately, while the coarser grains fall to the bottom, leaving the liquid quite clear after standing a minute or two. When the alcohol clears up in this way, it is an indication that it is not saturated, and an additional quantity of sugar should be shaken up with it. This is repeated until the mixture remains cloudy for at least two or three minutes after adding finely powdered sugar. When this takes place, the liquid is saturated, and a further addition of sugar will not increase its density. By operating in this way, we may obtain saturated alcohol in 10 or 15 minutes, while, by hanging sugar crystals in the solution, I have not been able to obtain the maximum of density in less than two days. When we are about to use the solution, it is advisable to shake it up a little while before using it, and, after the sugar has been deposited, to take the alcohometric degree. This degree cannot be higher than that shown by previous observations, but it may become lower, if the bottle is not tightly closed, as the alcohol evaporates, leaving a residuum containing a greater proportion of water. There is no use in placing chloride of calcium tubes on the cork of the bottle in the vain hope that, in this way, water may be absorbed from the atmosphere. The lowering of the alcoholic degree takes place principally by the evaporation of the spirit, and nothing can replace tight stoppers. The tin cans, with screw caps, that are used for refined petroleum, are well adapted for holding methylic alcohol, and of these Coleman's *elevated swinging cans* are the most convenient for holding our standard solution.

When on testing the saturated alcoholic solution, we find that the degree is lower than required, we may raise it by adding more alcohol. The addition of alcohol causes the precipitation of a part of the sugar in solution, so that the quantity of alcohol to be added should be somewhat less than the quantity to be added to a mixture of alcohol and water, of which we desire to raise the degree by the addition of stronger alcohol.

If we have a certain volume, V , of alcohol and water, whose alcohometric degree is d , and we wish to raise this degree to D , with strong alcohol of degree Δ , if the volume of the latter alcohol to be added is call x , we shall have—

* Since this paper was written, I have found the following interesting coincidence:—The alcohometric degree $83\frac{1}{4}$ corresponds to a specific gravity of 0.83315 (Prof. MacCulloch's tables for Traill's alcohometer). On consulting Dr. Ure's table for methylic alcohol (*Phil. Mag.* [3], xix., 51), it may be seen that specific gravity 0.8531 corresponds to 87 per cent of methylic alcohol. Now, in the first part of this paper, I have said that ethylic alcohol of 87 per cent, saturated with sugar, gives the results which agree most closely with those of the optical saccharometer.

† I usually take 0.8 x .

$$Vd + x\Delta = (V + x) D,$$

$$\text{whence} \quad x = \frac{V(D-d)}{\Delta-D}$$

If we have 1000 c.c. of alcohol at 81 , and we wish to raise the degree to $83\frac{1}{4}$, with alcohol of 92 per cent, then $d=81$, $D=83\frac{1}{4}$, $V=1000$ and $\Delta=92$, and the volume of alcohol of 92 to be added, is—

$$x = \frac{1000 \times 2.5}{8.5} = 294.1 \text{ c.c.}$$

If the addition of alcohol has been too great, we may diminish the degree by adding water very cautiously, and stirring up the mixture with an excess of sugar. To obtain the quantity of water, we may use the above formula but we must note that $\Delta=0$, and, as both numerator and denominator have become negative quantities, we may change the signs, and we shall have:—

$$x = \frac{V(d-D)}{D}$$

These particulars have been entered into because of the great importance of obtaining saturated solutions which do not deviate too widely from 77.1° of the alcohometer. Too much attention cannot be given to this, as otherwise it is not possible to obtain correct results.

Next in importance, is the weight of commercial sugar to be taken for 100 c.c. of the methylic solution saturated with sugar. In the process proposed by M. Dumas, 50 grms. of sugar are stirred with 100 c.c. of his standard solution. At the outset of the experiments on this process, I found that one obvious cause of error was the imperfect solubility of the impurities of commercial sugars in this standard solution. I was led then to grind the sugar down in a mortar as thoroughly as possible, so as to break up all the lumps and crush the large crystals, to allow everything to dissolve that would. I used for this quite a heavy pestle, of a size quite disproportionate to that of the mortar. By this thorough crushing much less sugar is used than M. Dumas proposed. It is an easy matter to determine the quantity of sugar to be used in a test. At first we may take an arbitrary quantity, and note the result, which may be corrected by the following considerations. The lowering of the alcohometric degree depends, as we have said, on the water and the soluble impurities present in the sugar. If we take a certain weight of sugar, say 45 grms., we may find by the alcohol process that the result is 91.5 per cent of sugar. If we test the same sugar by the optical saccharometer and find 93 per cent of sugar, we know that the alcohol process has given us too low a result, and this because the solution was too dense. Our result shows in the sugar:— $100-91.5=8.5$ of impurities and water, while we ought to have had $100-93=7$. We must then, to obtain 93, take a weight equal to—

$$\frac{45 \times 7}{8.5} = 37.05 \text{ grms.}$$

(To be continued.)

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 4, July 28, 1879.

Researches on the Refraction of Obscure Heat.—P. Desains.—In the case of a flint-glass prism acting upon the rays of incandescent platinum the spectrum is symmetrical with respect to the ordinate of the maximum as far as about 1° from this ordinate, either to the right

or the left. When this limit is passed the ordinates representing thermic intensities decrease rather more rapidly on the side of the luminous part of the spectrum than of the other. If crown glass prisms are used this symmetry no longer occurs.

Chloral Hydrate.—A. Wurtz.—When vapour of water and vapour of anhydrous chloral meet under such circumstances that no condensation takes place the mixture does not occasion the slightest rise of temperature. Hence the author concludes that no combination takes place and that the dissociation of the vapour of chloral hydrate is complete not merely at 100°, but under a reduced pressure even at 61°.

Observations on MM. Nobel and Abel's Memoir on Explosives.—M. Berthelot.—The author criticises the chemical equations representing the destruction of the explosive bodies, the measure of the heat disengaged, and the presence of potassium hyposulphite among the resulting products.

Researches on Samarium, the Radicle of a New Earth extracted from Samarskite.—Lecoq de Boisbaudran.—The spectrum of this metal is distinguished from that of the decipium of M. Delafontaine by the absence of the intense ray 478, and by the presence of the strong blue bands 480 and 463·5, the former of which covers the decipium ray 478, and lastly by the constant presence of the very strong band 400·75. The purification of the oxide of this metal is so tedious that the author has not yet obtained a quantity sufficient for the determination of its chemical properties.

Report on the Experimental Researches of M. Stanislas Meunier Relative to Meteoric Nickeliferous Irons and to the Native Iron Carbides of Greenland.—MM. Frey, H. Sainte-Claire Deville, Des Cloizeaux, Debray, and Daubrée.—The facts advanced justify the name of "veiny" (*filoniennes*) applied to concreted syssiderites. They even enable us to foresee that this name may be applied not merely to the numerous holosiderites remarkable for their crystalline condition, but even to the greater part of the metallic granulations of the sporadosiderites. The observations concerning the masses of Greenland compel us to admit that the internal layers of our globe, whence these masses are derived, are deeply impressed with a veiny character, all the rocks there being impregnated with concretions the result of reactions analogous to those which in ancient geological epochs gave rise to the formation of deposits of tin and of oligist. If we thus admit the analogy between meteorites and the deeper masses of our globe we recognise that the geological part of chlorine made known by Ch. Sainte-Claire Deville in his *Etudes sur les Volcans* is strikingly corroborated.

Action of Light upon Batteries.—H. Pellat.—A Daniell element whose copper is very clean is quite insensible to light. It is not the same if the copper is modified by oxidation or by the formation of a salt on its surface. Two Daniell elements were prepared as standards of electromotoric force, the sulphates being contained in two concentric glass vessels. These elements, perfectly transparent, were kept for five months; the zinc was not affected but the copper became coated with verdigris. Nevertheless, the elements retained their original force when the measurement was effected in the shade, but on exposure to the sun it was diminished by one-fortieth of its value. This variation ceased when the sun's rays were intercepted by a screen. This phenomenon is not due to a rise of heat, for the immersion of the battery in water at 50° produced no sensible effect, and a red glass which transmitted half the solar thermic radiations acted like an opaque screen. The luminous action renders the copper less positive.

Refrigerating Power of the Air at High Pressures.—A. Witz.—The law of Dulong and Petit represents exactly the facts up to a pressure of 1200 m.m.; at 1600 m.m.

the exponent of pressure requires to be nearly doubled; beyond that point it returns towards its first value, which it reaches about 6400 m.m.

Distillation of a Heterogeneous Liquid.—L. Troost.—In the distillation of a mixture of chloral hydrate and chloroform it is the water which distils with the chloroform as long as this exists in sufficient quantity. Nevertheless at 61° the maximum tension of the watery vapour is inferior by 50·2 m.m. to that of anhydrous chloral. If the anhydrous chloral remains in the retort with the chloroform though its vapour-tension is greater than that of water it is retained by a true chemical action. MM. Engel and Moitessier believe that the vapour which distils with chloroform at 61° contains at once water and anhydrous chloral. This is not the case.

Determination of Organic Matter in Natural Waters.—G. Lechartier.—The author, after having criticised the process of Prof. Frankland and shown that the evaporation with sulphurous acid occasions a loss of nitrogen, proposes to evaporate with magnesia, thus expelling all ammonia present in ammoniacal salts, and then to determine the total nitrogen present. A separate determination of nitric nitrogen gives the organic nitrogen as difference.

Thermo-chemical Study of Dissolved Alkaline Sulphides.—P. Sabatier.—The reaction of an alkali upon the hydrosulphate of a sulphide gives rise to a very sensible liberation of heat, the sign of the formation, in concentrated liquids of a certain dose of neutral sulphide.

Dissociation of Ammonium Hydrosulphate.—MM. Engel and Moitessier.—A reply to M. Isambert. A mixture of hydrogen sulphide and of ammonia in the cold is entirely absorbed by water. If the solution is heated the hydrogen sulphide escapes first, and is accompanied by but little ammonia.

Calcination of the Dregs of Beet-roots.—C. Vincent.—The author maintains, in opposition to MM. Du villier and Buisine, that his process is adequate to the preparation of considerable quantities of hydrochlorate of dimethyl-amine in a state of purity at least equalling that of the present commercial product.

Detection of Medicinal and Poisonous Substances in the Saliva.—A. Gabriel Pouchet.—In the case of patients suffering from lead poisoning lead was easily detected in the saliva. No arsenic could be detected in the case of diabetics under an arsenical treatment, nor could any trace of sugar be found. In *Morbus Brightii* albumen was present in the saliva.

The Soils of Dombes.—M. Nivet.—Remarkably poor soils which are kept under water for two seasons and are planted with oats in the third.

Experiments on the Production of Milk.—M. Lamj.—The author investigates the comparative influence of milking twice or thrice daily. He considers the latter system, other conditions being the same, more favourable to the production of butter.

No. 5, August 4, 1879.

Remarks on M. Wurtz's Note on Chloral Hydrate.—M. Berthelot.—The author points out that the method pursued by M. Wurtz in his experiments is not applicable when the amount of heat liberated is inconsiderable. The heat lost by radiation, contact, or conductivity is much greater than the quantity sought to be detected.

Normal Thermic Spectrum of the Sun, and of the Incandescent Platinum Lamp (Bourbouze Lamp).—M. Mouton.—As regards the solar spectrum the radiation of maximum intensity is very far from the point where Herschel's observations seemed to place it. It is between D and E, at the wave-length 0·56 μ . The violet and ultra-violet radiations present relatively considerable thermic intensities. The ultra-red portion is marked by four large known bands, the middle of which coincides

with the wave-lengths 0.85μ , 0.985μ , 1.23μ , and 1.48μ . The spectrum is extinct at 1.98μ much sooner than that of the Bourbouze lamp.

Vibrations on the Surface of Liquids.—A mathematical paper, not capable of useful abstraction.

On the Magnet.—M. Trève.—An indefinite rectilinear current acting successively and in the manner of friction upon a bar of soft iron slightly magnetised can cause it to lose completely or to resume in part its original magnetisation.

Distillation of Liquids under the Influence of Static Electricity.—D. Gernez.—Under the influence of static electricity there is a passage of liquids from the positive to the negative region, and this distillation results in nowise from the unequal heating of the two liquid strata traversed by the electricity. The quantity of liquid transported is proportional to the quantity of electricity brought into play, and does not depend on the extent of the free surface of the liquid.

Non-existence of a Soluble Alcoholic Ferment.—D. Cochin.—The author's experiments tend to demonstrate that a soluble alcohol ferment does not exist, and that vegetation is a direct and immediate consequence of the life of the yeast-cells.

Solid Hydride of Cyanogen.—H. Lescœur and A. Rigaut.—This compound may be formed by adding a trace of melted potassium cyanide to pure anhydrous hydrocyanic acid. It forms crystals, which in percentage composition agree with hydrocyanic acid. They dissolve readily in acids and appear feebly basic.

Synthetic Methyl-propyl-carbinol.—J. A. Le Bel.—The author proposes to use low forms of vegetation, such as *Penicillium glaucum*, for the purpose of separating isomeric bodies.

Colouring Matter of Palmella cruenta.—T. L. Phipson.—The *Palmella cruenta* is a small alga much resembling clotted blood. Its colouring matter, *palmellin*, closely resembles hæmoglobin. It is, like the latter, insoluble in alcohol, ether, benzol, sulphide of carbon, &c., but it dissolves in water. It is, further, dichroic; it is composed of red matter united to an albuminoid substance, and is coagulated by alcohol, acetic acid, and by the application of heat. Like the colouring matter of blood it produces absorption-bands in the yellow part of the spectrum, though not exactly in the same position. *Palmellin* easily enters into putrefaction, giving off a strong ammoniacal odour, and, to complete its analogies with the colouring matter of blood, it contains iron.

MISCELLANEOUS.

British Association for the Advancement of Science.—The following are the names of the Officers and Committee of Section B (Chemical Science Section) of the Sheffield Meeting of the British Association:—

President.—Prof. Dewar, M.A., F.R.S. L. and E.

Vice-Presidents.—Prof. Abel, F.R.S.; Dr. Longstaff; Lowthian Bell, M.P.; W. Crookes, F.R.S.; Dr. J. H. Gladstone, F.R.S.; Dr. Gilbert, Ph.D., F.R.S.; A. Vernon Harcourt, M.A., F.R.S.; Prof. Odling, M.B., F.R.S.; Dr. Ronalds, F.R.S.E.; H. Clifton Sorby, F.R.S.; Prof. A. W. Williamson, F.R.S.

Secretaries.—W. Chandler Roberts, F.R.S.; J. Millar Thomson (Recorder); H. S. Bell, F.C.S.

Committee.—A. H. Allen, G. Ansdell, Prof. Attfield, Dr. Aug. Bernthsen, Dr. Campbell Brown, P. Braham, J. Y. Buchanan, F.R.S.E.; P. H. Carpenter, M.A.; M. P. de Clermont, J. H. Collins, W. Fairley, F.R.S.E.; A. Fletcher, Col. J. C. Gamble, W. Gillingham, G. Gladstone, W. N. Hartley, F.R.S.E.; A. K. Huntington, F. M. Jennings, Taiso Masaki, Prof. McLeod, E. Macadam, Dr. Morgan, Prof. Rowney, G. F. Schacht, M. R. D. Silva, Peter Spence, E. C. Stanford, W. Stoddard, W. Thomson,

F.R.S.E.; R. C. Tichborne, Dr. Wm. Wallace, A. Wanklyn, R. Warrington, W. Weldon, F.R.S.E.; E. J. Williams.

The papers brought before the Section were as follows:—

W. Chandler Roberts, F.R.S.—Report of Committee on the Chemistry of some of the lesser-known Alkaloids.

Walter Weldon, F.R.S.E.—On some relations between the numbers expressing the Atomic Weights of the Elements.

M. R. D. Silva.—On the Synthesis of Diphenyl-propyl.

F. A. Abel, F.R.S.—Recent Researches in Explosive Agents.

Prof. Dewar, F.R.S.—On Vapour-Densities.

P. Braham.—To Describe a large Crystal of Mercury Sulphate.

Henry S. Bell, F.C.S.—On the Manufacture of Crucible Steel.

Thomas Blair.—On the Separation of Iron and Phosphorus, especially with reference to the Manufacture of Steel.

John Hollway.—A New Process in Metallurgy.

A. H. Allen, F.C.S.—A Lecture Experiment in Illustration of the Hollway Process of Smelting Sulphide Ores.

Andrew French.—On Lead Fume, with a description of a New Process of Fume Condensing.

Prof. Odling, F.R.S.—On the Constitution of Aluminic Compounds.

A. H. Allen.—On the Presence of Nitrogen in Steel.

A. Vernon Harcourt, F.R.S.—Colour Tests for Phosphorus and Sulphur in Iron and Steel.

W. Chandler Roberts, F.R.S.—To Exhibit some Experiments with Hughes's Voltaic Induction Balance.

J. T. Brown.—Historical Sketch of the various Vapour-Density Methods.

Prof. Wanklyn.—Note on Certain Vapour-Densities.

Prof. Wanklyn.—Note on Isocyan-propionic Acid.

G. Ansdell.—Physical Constants of Liquid Acetylene and Hydrochloric Acid.

M. de Clermont.—The Action of Ammoniacal Salts on Metallic Sulphides.

W. Ivison Macadam.—On the Chemical Composition of a Nodule of Ozokerite found at Kinghorn-ness.

Thos. Andrews.—On some Curious Concretion Balls derived from a Colliery Mineral Water.

Dr. Gilbert, F.R.S.—On some points in connection with Agricultural Chemistry.

T. S. Humpidge, B.Sc.—On the rare Metals of the Yttrium Group.

Prof. Dewar, F.R.S.—On the Synthesis of Hydrocyanic Acid.

Prof. Dewar, F.R.S.—On the Amount of Nitrous Acid produced in Electric Illumination.

Prof. Dewar, F.R.S.—On the Kinoline Bases.

John M. Thomson.—An account of some Recent Experiments on Supersaturated Solutions.

J. Norman Lockyer, F.R.S.—Notes of some Recent Spectral Observations.

A. H. Allen.—Notes on Petroleum Spirit or Benzoline.

A. Vernon Harcourt, F.R.S.—On the Illuminative Value of a Mixture of Hydrogen.

G. T. Hazelhurst.—On a New Form of Condenser.

W. Thomson, F.R.S.E.—Notes on a Sample of Fuller's Earth found in an old Fullonica recently excavated at Pompeii.

W. H. Watson.—On the Detection of Milk Adulteration.

Dr. Phipson.—Chemical Researches on the *Palmella cruenta*.

Philip Braham.—Description of a Glass Burette for Collecting, Measuring, and Discharging Gas over Mercury.

TO CORRESPONDENTS.

J. M. Caven.—Metallic zinc is prepared by distillation and rapid condensation. It can be purchased cheaply.
Blue Blood.—In *Reimann's Farber Zeitung*.

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THE SUPPOSED COMPOUND NATURE OF THE ELEMENTS.

By J. NORMAN LOCKYER, F.R.S., &c.

CONTINUING my researches into the nature of the so-called elements, I have found that when carefully distilled metallic sodium was condensed in a capillary tube, placed in a retort, and heated in a Sprengel vacuum it gave off twenty times its volume of hydrogen. Phosphorus, carefully dried and submitted to the same treatment, gave off 70 volumes of a gas which appeared to consist chiefly of hydrogen. Although it gave some of the lines of phosphorus it was not PH_3 , as it had no action on solution of cupric sulphate. A specimen of magnesium carefully purified by Messrs. Johnson and Matthey, gave me a magnificent series of coloured phenomena. The hydrogen lines first appeared, then the D line—not the sodium line be it understood, for the green line was absent—and, lastly, the green line of magnesium (b), and then, as the temperature was increased, mixtures of all these lines, with the blue line, the D line being always the most brilliant. In this experiment only 2 volumes of hydrogen were collected. From gallium and arsenic no gas of any kind was obtained. From sulphur and some of its compounds sulphurous anhydride was always obtained. From indium hydrogen was given off *in vacuo* before heating, while from lithium no less than 100 volumes of hydrogen were given off. The conditions of the experiments were always the same, the only variable being the substance itself.

A LECTURE EXPERIMENT IN ILLUSTRATION OF THE HOLLWAY PROCESS OF SMELTING SULPHIDE ORES.*

By ALFRED H. ALLEN, F.I.C., F.C.S.

By causing oxygen gas to bubble through molten antimony sulphide contained in a V-shaped piece of combustion-tube, combustion takes place with such rise of temperature as to soften the glass, while a sublimate is obtained of antimonious iodide, and sulphurous acid gas is evolved. The sublimate is collected in an empty globe, and the sulphurous acid is absorbed by passing it into a large vessel containing lumps of wood-charcoal. At the conclusion of the experiment the contents of the combustion-tube may be poured out, when a button of metallic antimony free from sulphur is obtained.

By passing oxygen over lumps of pyrites contained in a heated combustion-tube, vivid combustion takes place, much free sulphur sublimes, and sulphurous acid gas is obtained and absorbed as before described.

NOTES ON PETROLEUM SPIRIT OR "BENZOLINE."

By ALFRED H. ALLEN, F.I.C., F.C.S.

THE application of the commercial names "benzoline" and "benzine" to the more volatile portion of petroleum

has led to great confusion between petroleum spirit and coal-tar naphtha, the most characteristic constituent of which is the hydrocarbon *benzene* or *benzol*.

Although presenting close general resemblances, the following characteristic differences exist between petroleum spirit and coal-tar naphtha. All the tests given have been carefully verified by the author on representative samples of petroleum spirit and coal-tar benzol.

Petroleum Spirit, "Benzoline" or "Benzine."

1. Consists of *heptane*, C_7H_{16} , and its homologues.
2. Heptane contains 84.0 per cent of carbon.
3. Commences to boil at 54° to 60° C.
4. Specific gravity at 15.5° C. about 0.69 to 0.72.
5. Smells of petroleum.
6. Dissolves iodine, forming a solution of a raspberry-red colour.
7. Does not sensibly dissolve coal-tar pitch, and is scarcely coloured by it, even on prolonged contact.
8. When shaken in the cold, with one-third of its volume of fused crystals of absolute carbolic acid, the latter remains undissolved, and forms a separate lower stratum.
9. Requires two volumes of absolute alcohol, or four or five volumes of methylated spirit of 0.828 sp. gr., for complete solution at the ordinary temperature.
10. Warmed with four measures of nitric acid of 1.45 sp. gr. the acid is coloured brown, but the spirit is little acted on, and forms an upper layer.

Coal-Tar Naphtha, or "Benzol."

1. Consists of *benzene*, C_6H_6 , and its homologues.
2. Benzene contains 92.3 per cent of carbon.
3. Commences to boil at about 80° C.
4. Specific gravity about 0.88.
5. Smells of coal-tar.
6. Dissolves iodine, forming a purple-red liquid of the tint of an aqueous solution of potassium permanganate.
7. Readily dissolves coal-tar pitch, forming a deep-brown solution.
8. Miscible with absolute carbolic acid in all proportions.
9. Miscible with absolute alcohol in all proportions. Forms a homogeneous liquid with an equal measure of methylated spirit of 0.828 sp. gr.
10. Completely miscible with four measures of nitric acid of 1.45 sp. gr., with great rise of temperature and production of dark brown colour. A portion of the nitrobenzene produced may separate as the liquid cools.

The greater number of the above tests are valueless when applied to mixtures of petroleum and coal-tar naphthas, but No. 10 is capable of giving quantitative results if the treatment with nitric acid be conducted in a small flask and an inverted condenser attached, to prevent loss of vapours. When action has nearly ceased, if the liquid be poured into a narrow graduated tube, the measure of the upper layer indicates with approximate accuracy the amount of petroleum spirit present. If the proportion of benzene is considerable, the nitrobenzene produced may not remain completely dissolved in the nitric acid, in which case it rises and forms a layer of a dark brown colour below the stratum of petroleum spirit. Nitrobenzene and petroleum spirit are readily miscible in the absence of nitric acid, but agitation with strong nitric acid dissolves out the nitrobenzene, a portion of which may rise and form an intermediate layer as above described. By fractional distillation, the author found that the

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

proportion of *heptane*, C_7H_{16} , present in commercial benzoline probably equalled, or even exceeded, that of all the other constituents.

ON THE DETECTION OF MILK ADULTERATION.*

By WILLIAM H. WATSON, F.C.S., &c.

FROM analyses of milk from various dairies, and by a comparison of the results obtained with circumstances existing as to the character and quantity of the food; nature of different cows; conditions and health of them at particular periods; and changes of the seasons of the year, the author concludes that cows' milk is subject to considerable variations in composition. He has found in many instances milk from well-fed healthy cows to contain as little as 10·5 per cent of total solids, and from 8·5 to 9 per cent of solids not fat. The results of other experimenters are compared, and it is then suggested that the present limits adopted by public analysts for genuine milk should be re-considered.

ON THE MANUFACTURE OF CRUCIBLE STEEL.*

By HENRY S. BELL, F.C.S., &c.

THE manufacture of crucible steel is one of the most important industries connected with the town of Sheffield, which boasts of not less than 120 firms engaged in the production of this material. Notwithstanding the enormous output of steel by the Bessemer and Siemens-Martin processes, this kind of steel is unrivalled for the manufacture of the finer varieties of cutlery and edged tools, &c. A brief outline of the process itself is as follows:—The most of the iron employed for this purpose is imported into this country in the shape of bars, from Sweden, where it has been smelted from very pure iron ores, in a blast-furnace, by the aid of charcoal, and subsequently puddled to free it from impurities.

The first operation to which it is subjected is that known as the cementation or converting process, the object of which is to combine a certain quantity of carbon with the iron; this operation is performed in a furnace of peculiar construction, where the iron and charcoal are packed together in air-tight chests or converting pots, subjected to a high temperature short of the fusing-point of iron, where it remains for a matter of three weeks.

After the conversion, when the pots are cold the bars are taken out and found to be covered with blisters; hence it is termed Blister Steel. In consequence of the various theories proposed to account for this peculiar formation, the writer was induced to make a series of investigations. For this purpose he was kindly furnished by Messrs. Seebohm and Dieckstaht, of the Dannemora Steel Works, with some samples of this blister steel, various portions of which he submitted to analysis, the results of which showed a marked increase of silicon where the blisters occurred, thus—

Sample	Description	Per cent Silicon.
1.	Blister 2 ins. in length contained	0.070
2.	Small blister contained	0.048
3.	" " " " "	0.056
1.	Unblistered portions contained	0.023
2.	" " " " "	0.021
3.	" " " " "	0.025

On inspecting one of these bars of blister steel, it is found that it has undergone both a physical and a chemical change.

* Abstract of a paper read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

The iron has now assumed a crystalline structure, and has chemically combined with a certain amount of carbon. This latter change commences on the exterior, and extends itself to the interior of the bar, if the process be continued sufficiently long, thus showing that carbonic oxide never penetrates into the centre of the bar, until the whole is converted into steel.

The writer is indebted to the kindness of the above-mentioned firm for a sample of bar iron, before and after conversion, in order to ascertain the exact chemical change that took place during the process. The following are the results obtained :—

	Before Conversion.	After Conversion.
Fe	99'471	98'603
C	0'352	1'250
Si	0'050	0'035
S	0'027	0'022
P	0'025	0'018
Mu	0'075	0'072
	100'000	100'000

The decrease in impurities appears greater than it really is, owing to the fact that the bar itself has increased in weight by the addition of carbon.

One remarkable fact is that, after the conversion of the iron, a quantity of the charcoal, in the converting pot, is found in a pulverised state, so as to be unfit for further use.

Some of this waste charcoal the writer has examined, and from one sample, by the aid of a magnet, he succeeded in extracting 5 to 6 per cent of iron scale and small pieces of steel. These on being treated with dilute hydrochloric acid, evolved considerable quantities of sulphuretted hydrogen; in one case he estimated the quantity of sulphur, and found it to contain as much as 1.25 per cent of this element. The blister steel thus produced, for the sake of convenience is divided into six different classes, viz.

Spring heat. Country heat.
Single shear heat. Double shear heat.
Steel through heat. Melting heat.

The steel is now broken up into small pieces and melted in crucibles, and cast into ingots. These are sent to the forge, where they are heated and rolled. In this part of the process the chief difficulty with which the tilter has to contend is the porous or "honey-combed" structure of the steel.

One of the characteristic features relied on by practical men as indicating the quality of a piece of steel is the appearance of its fracture; but this is by no means an infallible test, as the fineness or coarseness of grain can be produced by mechanical treatment or chemical means.

The characteristic property possessed by steel is its capability of being hardened and tempered. The temper of cast steel may be said to range from 0.75 to 1.50 per cent carbon. The temper of steel is an important question in connection with the purpose for which it is required: thus a steel containing 1.50 per cent of carbon is the class employed for razors; 1.25 per cent is that known as "tool temper;" steel containing 1.00 per cent carbon is termed "chisel steel," and this temper is extensively used in the arts.

The favourite marks of Swedish non-employed in the manufacture of this kind of steel, are those obtained from Dannemora, the most noted of which are the 00=Double Bullet,—

$$G_L = GL, \text{ and } (\widehat{L}) = \text{Hoop } L.$$

The most important of the elements which affect the quality and mechanical properties of steel are the following :—

Carbon, Silicon, Sulphur, Phosphorus, and Manganese.
Carbon, by its direct combination with iron, is essentially

the steel-forming element, and greatly increases the hardness and tensile strength of the metal. The maximum quantity of carbon capable of being taken up by iron is 6·5 per cent to 7·00 per cent. This high percentage of carbon is only attained, as in the case of rich ferromanganese, containing as much as from 85 to 86 per cent of manganese.

Silicon.—The action of this element on steel is to produce both red- and cold-shortness, especially in high made steels. Under certain conditions, it is capable of imparting hardness without brittleness. The presence of this element also tends to favour a solid casting, and prevent the formation of a honey-combed structure.

Sulphur in steel, as is well known, produces "red-shortness," and has also a tendency to prevent the chemical combination of iron with carbon, and also to displace it when in combination.

Phosphorus produces cold-shortness and brittleness, but the detrimental influence of this element, when present only in small quantities, can be partially neutralised, providing the percentage of carbon is very low.

Manganese is a valuable ally of the steel melter, and serves to correct the evil effects produced by the presence of sulphur, oxygen, &c.; and when in the state of an oxide serves to eliminate a large percentage of the silicon.

Manganese is generally introduced into the steel in the form of "Spiegeleisen," an alloy of iron, carbon, and manganese, generally containing about 10 per cent of the latter element.

Other metals have been employed to replace carbon, such as tungsten, chromium, and titanium; these impart great hardness and fineness to the texture of steel.

For a considerable amount of practical information given in his paper, but necessarily omitted from this abstract, the writer is indebted to a valuable essay written some years ago on this subject by Henry Seeborn, of the firm of Messrs. Seeborn and Dieckstahl.

This paper is not intended to give any additional information to the practical steel makers of Sheffield, as to the manufacture of steel, or to offer any criticisms or advice in the matter; its object is simply to give an outline of the manufacture as it is still carried on in this town, with the hope that it may prove interesting to many of those who have come from a distance to attend the visit of the British Association, and who are unacquainted with the process which has caused Sheffield to become the great manufacturing steel centre in this country.

ON SOME CURIOUS CONCRETION BALLS DERIVED FROM A COLLIERY MINERAL WATER.*

By THOMAS ANDREWS, F.C.S.

THE water on which these observations were made was collected from the "sump" of the Wortley Silkstone Colliery, a small pit situated near the "Bassett" or "outcrop" of the great Silkstone seam of coal; the samples being obtained during typical dry and rainy seasons. The water had percolated from the surface a distance of 35 yards, through strata, as indicated on the accompanying table.

The bottom layer in which the water lodged was the Silkstone seam of coal, here some 5 feet in thickness.

One noticeable feature of this water is, that it always gives an acid reaction with blue litmus paper.

Several analyses of this water made at various times indicate that the chief mineral constituents of the water are, iron—calcium, magnesium, in the form of sulphates.

This water when heated quickly throws down a copious ochreous deposit. The deposit found in the engine boilers

after having used the water in them for steam purposes was of the composition given below.

The boiler residue from which this sample was taken consisted of an incrustation about one inch thick, which had adhered to the bottom of the boiler.

The incrustation was of a light reddish yellow colour in the bulk, it was very hard and tough, and not easily broken in pieces.

The iron work in connection with this colliery engine and boilers, in any way exposed to the action of either the acid water itself, or the steam generated from it, becomes corroded and partially dissolved. The most effectual remedy against this corrosive action and deposit, is that described in my letter to the CHEMICAL NEWS, vol. xxxv., p. 252.

Analysis of Boiler Deposit, from Wortley Silkstone Colliery Boilers, September 15, 1875.

Moisture	6·85	per cent.
Combined water, organic matter, &c. ..	5·80	"
Silicious matter	1·80	"
Peroxide of iron and alumina containing } phosphoric acid 0·76 per cent }	6·10	"
Sulphate of lime	78·55	"
Magnesia	0·65	"
	99·75	"

Some curious balls of mineral matter are occasionally found in the feed tank of the colliery boilers, which are supplied with this water. The water is pumped up from the engine pond into a cylindrical feed tank, and is there heated by the exhaust steam from the engine playing on its surface (not blowing through it). The water in this feed tank has an average temperature of 164° F.

It sometimes happens that during the short space of even two or three weeks, great numbers of these balls are formed, varying in size from about 3½ inches in diameter to ½ inch diameter, and in weight from about one and a half pounds to a quarter of an ounce.

The author has many of these in his possession. They are perfectly hard and compact when taken from the tank, and are no doubt formed from the deposit thrown down when the mineral water is heated.

The action of steam playing on the surface of the water probably causes circular eddies, and when a nucleus has thus once been formed, it is easy to conceive of the gradual formation and consolidation of these balls.

The author suggests that the conditions of formation of natural nodules of iron ore, pyrolusite, &c., may be similar to those observed by him in the foregoing cases.

The following is the analysis of the balls formed in the feed tank, from which it will be seen that they are quite different in composition to the residue deposited in the boilers, owing probably to the difference in the temperature between the feed tank and the boiler.

Analysis of a Concretion Ball, found in the Engine Feed Tank, Wortley Silkstone Colliery.

Moisture	2·30	per cent.
Loss on ignition, organic matter, &c., } contains matters extracted by ether, } 5·8 per cent	24·40	"
Silica	1·80	"
Peroxide of iron	62·86	"
Alumina	5·43	"
Phosphoric acid	2·81	"
Lime	0·40	"
Magnesia	trace	"
	100·00	"

They also differ in composition from concretions deposited at the pit bottom in the cold, which show only 45·71 per cent of Fe₂O₃.

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

Observations made to ascertain the Temperature at which the Deposit and Turbidity take place.

Temperature at which Turbidity Commences.
Dry Season. Wet Season.
Average of four observations .. 147° F. 160.5° F.

The action of this mineral water is destructive to all iron work with which it comes in contact. The amount of iron dissolved by samples of the water in the cold being as follows:—

July, 1868.—7000 fluid grains = 1 lb. of the water dissolved 1.85 grain of iron during one month.

Feb., 1876.—3500 fluid grains = $\frac{1}{4}$ lb. of the water (collected during a dry season) dissolved 2.91 grains of iron in eight months.

July, 1876.—3500 fluid grains = $\frac{1}{4}$ lb. of the water (collected during a dry season) dissolved 4.73 grains of iron in eight months.

Reaction of the water with litmus at the conclusion of these experiments only faintly acid.

Quantities of iron-pyrites are found in the coal strata of this neighbourhood, and account for the large quantities of sulphates often found in these colliery waters.

The water during flood seasons required (as an average of five determinations) an addition of 10.48 grains of anhydrous Na_2O per gallon, before an alkaline reaction was obtainable, and during dry seasons (as an average of three determinations) an addition of 17.35 grains of anhydrous Na_2O per gallon. This amount of alkali does not all correspond to free acid, as the sulphate of iron would also neutralise soda.

Determinations of the Total Inorganic Constituents were made at the dates and with the results as below. Results in grains per gallon.

Date.	Total amount of Inorganic Matters.
June 20, 1865.. ..	56.20
October 12, 1867	66.60
July 27, 1868	67.60 (Very dry season)
June 25, 1874.. ..	133.70
June 27, 1874.. ..	198.80 (Very dry season)

The sulphates, a very important element in the composition of this water, were determined as per following results. An average of six estimations of the total sulphates (results calculated as SO_3), extending from 1865 to 1876, made during—

Dry seasons—giving 115.41 grains per gallon of SO_3 .

An average of eight estimations, extending over the same period of time, but made during—

Rainy seasons—gave 67.62 grains per gallon of SO_3 .

A fact worthy of notice in course of these analyses is the steady and large increase in the amount of the sulphates, from the commencement of these observations in 1865 to 1876.

The same result is also noticeable on reference to the total amounts of inorganic matter, which show a great increase in quantity during the latter part of the time.

Now why this increase in the total sulphates and total matters in solution should take place it is not easy to say. It may be owing to the fact of the increased length and area of the workings in the colliery, as undoubtedly there is more bulk of water to contend with now than formerly on this account; but why the mineral matter in the water should have increased owing simply to this increase in quantity is not at first sight very clear.

It may be possible that the largely increased number, area, and length of air-ways has a tendency to expose the water for a longer time to oxidising influences, and thus add to the percentage of sulphates, and this increased facility for oxidation no doubt also induces more rapid solution of the strata as the water slowly permeates through it.

The author hopes these few imperfect observations may not prove altogether uninteresting to those who take pleasure in the study of mineral waters.

ON RADIANT MATTER.*

By WILLIAM CROOKES, F.R.S.

(Continued from page 93.)

Radiant Matter proceeds in straight lines.

THE Radiant Matter whose impact on the glass causes an evolution of light, absolutely refuses to turn a corner. Here is a V-shaped tube (Fig. 6), a pole being at each ex-

FIG. 6.



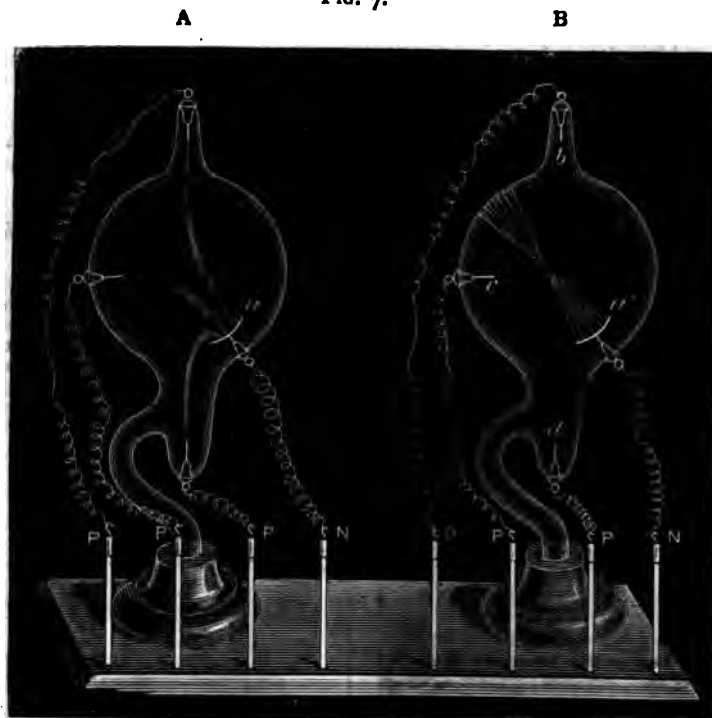
tremity. The pole at the right side (α) being negative, you see that the whole of the right arm is flooded with green light, but at the bottom it stops sharply and will not turn the corner to get into the left side. When I reverse the current and make the left pole negative, the green changes to the left side, always following the negative pole and leaving the positive side with scarcely any luminosity.

In the ordinary phenomena exhibited by vacuum tubes—phenomena with which we are all familiar—it is customary, in order to bring out the striking contrasts of colour, to bend the tubes into very elaborate designs. The luminosity caused by the phosphorescence of the residual gas follows all the convolutions into which skilful glass-blowers can manage to twist the glass. The negative pole being at one end and the positive pole at the other, the luminous phenomena seem to depend more on the positive than on the negative at the ordinary exhaustion hitherto used to get the best phenomena of vacuum tubes. But at a very high exhaustion the phenomena noticed in ordinary vacuum tubes when the induction spark passes through them—an appearance of cloudy luminosity and of stratifications—disappear entirely. No cloud or fog whatever is seen in the body of the tube, and with such a vacuum as I am working with in these experiments, the only light observed is that from the phosphorescent surface of the glass. I have here two bulbs (Fig. 7), alike in shape and position of poles, the only difference being that one is at an exhaustion equal to a few millimetres of mercury—such a moderate exhaustion as will give the ordinary luminous phenomena—whilst the other is exhausted to about the millionth of an atmosphere. I will first connect the mode-

* A Lecture delivered to the British Association for the Advancement of Science, at Sheffield, Friday, August 22, 1879.

exhausted bulb (A) with the induction-coil, and re- upper end of the tube is another terminal, *d*. The induc-
the pole at one side (*a*) always negative, I will put tion-coil is connected so that the hemi-cylinder is nega-

FIG. 7.



live wire successively to the other poles with which
lb is furnished. You see that as I change the
a of the positive pole, the line of violet light joining
poles changes, the electric current always choosing
shortest path between the two poles, and moving about
b as I alter the position of the wires.

, then, is the kind of phenomenon we get in ordinary
stions. I will now try the same experiment with a bulb
it is very highly exhausted, and as before, will make
e pole (*a'*) the negative, the top pole (*b*) being positive.
how widely different is the appearance from that
by the last bulb. The negative pole is in the
f a shallow cup. The molecular rays from the cup
n the centre of the bulb, and thence diverging, fall on
posite side and produce a circular patch of green
rescent light. As I turn the bulb round you will
ble to see the green patch on the glass. Now observe,
ve the positive wire from the top, and connect it
he side pole (*c*). The green patch from the divergent
ra focus is there still. I now make the lowest pole
sitive, and the green patch remains where it was at
nchanged in position or intensity.

have here another property of Radiant Matter. In
r vacuum the position of the positive pole is of every
ance, whilst in a high vacuum the position of the
a pole scarcely matters at all; the phenomena seem
nd entirely on the negative pole. If the negative
ints in the direction of the positive, all very well,
he negative pole is entirely in the opposite direction
little consequence: the Radiant Matter darts all
ne in a straight line from the negative.

instead of a flat disk, a hemi-cylinder is used for the
re pole, the Matter still radiates normal to its sur-
The tube before you (Fig. 8) illustrates this pro-

It contains, as a negative pole, a hemi-cylinder (*a*)
shed aluminium. This is connected with a fine
wire, *b*, ending at the platinum terminal, *c*. At the

FIG. 8.



tive and the upper pole positive, and when exhausted to a
sufficient extent the projection of the molecular rays to a
focus is very beautifully shown. The rays of Matter being

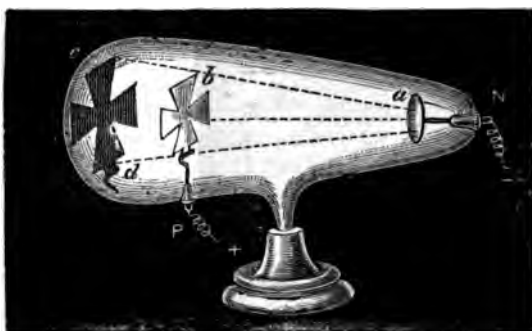
driven from the hemi-cylinder in a direction normal to its surface, come to a focus and then diverge, tracing their path in brilliant green phosphorescence on the surface of the glass.

Instead of receiving the molecular rays on the glass, I will show you another tube in which the focus falls on a phosphorescent screen. See how brilliantly the lines of discharge shine out, and how intensely the focal point is illuminated, lighting up the table.

Radiant Matter when intercepted by solid matter casts a shadow.

Radiant Matter comes from the pole in straight lines, and does not merely permeate all parts of the tube and fill it with light, as would be the case were the exhaustion less good. Where there is nothing in the way the rays strike the screen and produce phosphorescence, and where solid matter intervenes they are obstructed by it, and a shadow is thrown on the screen. In this pear-shaped bulb (Fig. 9) the negative pole (a) is at the pointed end.

FIG. 9.



In the middle is a cross (b) cut out of sheet aluminium, so that the rays from the negative pole projected along the tube will be partly intercepted by the aluminium cross, and will project an image of it on the hemispherical end of the tube which is phosphorescent. I turn on the coil, and you will all see the black shadow of the cross on the luminous end of the bulb (c, d). Now, the Radiant Matter from the negative pole has been passing by the side of the aluminium cross to produce the shadow; the glass has been hammered and bombarded till it is appreciably warm, and at the same time another effect has been produced on the glass—its sensibility has been deadened. The glass has got tired, if I may use the expression, by the enforced phosphorescence. A change has been produced by this molecular bombardment which will prevent the glass from responding easily to additional excitement; but the part that the shadow has fallen on is not tired—it has not been phosphorescing at all and is perfectly fresh; therefore if I throw down this cross,—I can easily do so by giving the apparatus a slight jerk, for it has been most ingeniously constructed with a hinge by Mr. G. mingham,—and so allow the rays from the negative pole to fall uninterruptedly on to the end of the bulb, you will suddenly see the black cross (c, d, Fig. 10) change to

FIG. 10.



a luminous one (e, f), because the background is now only capable of faintly phosphorescing, whilst the part which had the black shadow on it retains its full phosphorescent power. The stencilled image of the luminous cross unfortunately soon dies out. After a period of rest the glass partly recovers its power of phosphorescing, but it is never so good as it was at first.

Here, therefore, is another important property of Radiant Matter. It is projected with great velocity from the negative pole, and not only strikes the glass in such a way as to cause it to vibrate and become temporarily luminous while the discharge is going on, but the molecules hammer away with sufficient energy to produce a permanent impression upon the glass.

Radiant Matter exerts strong mechanical action where it strikes.

We have seen, from the sharpness of the molecular shadows, that Radiant Matter is arrested by solid matter placed in its path. If this solid body is easily moved the impact of the molecules will reveal itself in strong mechanical action. Mr. G. mingham has constructed for me an ingenious piece of apparatus which when placed in the electric lantern will render this mechanical action visible to all present. It consists of a highly exhausted glass tube (Fig. 11), having a little glass railway running along

FIG. 11.



it from one end to the other. The axle of a small wheel revolves on the rails, the spokes of the wheel carrying mica paddles. At each end of the tube, and rather above the centre, is an aluminium pole, so that whichever pole is made negative the stream of Radiant Matter darts from it along the tube, and striking the upper vanes of the little paddle-wheel causes it to turn round and travel along the railway. By reversing the poles I can arrest the wheel and send it the reverse way, and if I gently incline the tube the force of impact is observed to be sufficient even to drive the wheel up-hill.

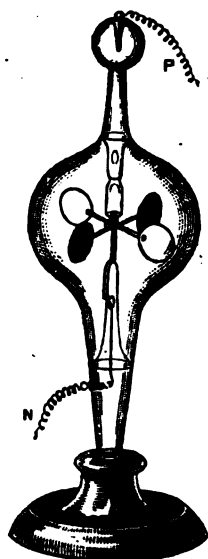
This experiment therefore shows that the molecular stream from the negative pole is able to move any light object in front of it.

The molecules being driven violently from the pole there should be a recoil of the pole from the molecules, and by arranging an apparatus so as to have the negative pole movable and the body receiving the impact of the Radiant Matter fixed, this recoil can be rendered sensible. In appearance the apparatus (Fig. 12) is not unlike an ordinary radiometer with aluminium disks for vanes, each disk coated on one side with a film of mica. The fly is supported by a hard steel instead of glass cup, and the needle point on which it works is connected by means of a wire with a platinum terminal sealed into the glass. At the top of the radiometer bulb a second terminal is sealed in. The radiometer therefore can be connected with an induction-coil, the movable fly being made the negative pole.

For these mechanical effects the exhaustion need not be so high as when phosphorescence is produced. The best pressure for this electrical radiometer is a little beyond that at which the dark space round the negative pole extends to the sides of the glass bulb. When the pressure is only a few millims. of mercury, on passing the induc-

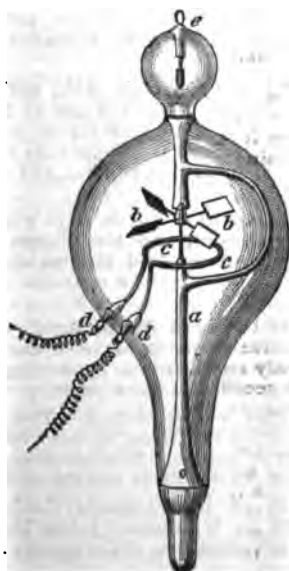
tion current a halo of velvety violet light forms on the metallic side of the vanes, the mica side remaining dark. As the pressure diminishes, a dark space is seen to separate the violet halo from the metal. At a pressure of

FIG. 12.



half a millim. this dark space extends to the glass, and rotation commences. On continuing the exhaustion the dark space further widens out and appears to flatten itself against the glass, when the rotation becomes very rapid.

FIG. 13.



Here is another piece of apparatus (Fig. 13) which illustrates the mechanical force of the Radiant Matter from the negative pole. A stem (a) carries a needle-point in which revolves a light mica fly (bb). The fly consists of four square vanes of thin clear mica, supported on light aluminium arms, and in the centre is a small glass cap which rests on the needle-point. The vanes are inclined

at an angle of 45° to the horizontal plane. Below the fly is a ring of fine platinum wire (cc), the ends of which pass through the glass at dd. An aluminium terminal (e) is sealed in at the top of the tube, and the whole is exhausted to a very high point.

By means of the electric lantern I project an image of the vanes of the screen. Wires from the induction-coil are attached, so that the platinum ring is made the negative pole, the aluminium wire (e) being positive. Instantly, owing to the projection of Radiant Matter from the platinum ring, the vanes rotate with extreme velocity. Thus far the apparatus has shown nothing more than the previous experiments have prepared us to expect; but observe what now happens. I disconnect the induction-coil altogether, and connect the two ends of the platinum wire with a small galvanic battery; this makes the ring cc red-hot, and under this influence you see that the vanes spin as fast as they did when the induction-coil was at work.

Here, then, is another most important fact. Radiant Matter in these high vacua is not only excited by the negative pole of an induction-coil, but a hot wire will set it in motion with force sufficient to drive round the sloping vanes.

(To be continued.)

NEW PROCESS FOR THE RAPID ESTIMATION OF PURE SUGAR IN RAW AND REFINED COMMERCIAL SUGARS.*

By P. CASAMAJOR.

(Continued from page 98.)

AFTER trying many experiments with solutions of different strengths, it was found that each solution required a different weight. For the saturated solution of 77.1° of the alcohometer, which is our standard solution, the weight is 39.6 grms. for 100 c.c. of the solution.

Instead of using 100 c.c., I have, for a long time, used only 50. To be able to use a cylinder in which this volume would give indications, I needed to use alcohometers of quite small diameter. These were made by Mr. H. Weinhausen; they are very well graduated and very satisfactory in every way.

For 50 c.c. of standard solution, the proper weight is half of the one for 100 c.c. = 19.8 grms. To show how this was obtained, the following table, No. 2. is given. In the first column are numbers designating the sugars that were tested; in the second column is the percentage of sugar as obtained by the optical saccharometer; in the third column is the result by the methylic alcohol process, when we take 19 grms. as the weight of the sugar tested; in the fourth column is the quantity that should have been taken, instead of 19 grms., to obtain the same result as by the optical saccharometer. These weights were obtained by the calculation already given.

TABLE No. 2.

Sugar.	Degree of saccharometer.	Result by methylic alcohol. Weight, 19 gr.	Weight to give results in 2nd col. gr.
1	91.6	91.9	19.7
2	84.8	85.36	19.66
3	91.7	92.1	19.96
4	81.7	82.46	19.8
5	88.8	89.1	19.7
6	96.3	96.6	20.6

From the results in the 4th column, the weight adopted was 19.8 gr.

By using this weight, with 50 c.c. of standard solution the following results were obtained in the case of 15 consecutive tests of raw and refined sugars:—

* Read before the American Chemical Society, June 5, 1879.

TABLE No. 3.

Designation of the sugar.	Percentage of pure sugar by saccharometer.	Ditto by methylic alcohol.	Difference.
Muscavado ..	82.4	82.3	-0.1
" ..	91.5	91.7	0.2
Refined ..	95.0	95.1	0.1
Molasses ..	86.5	87.2	0.7
Centrifugal ..	96.5	96.2	-0.3
Refined ..	96.3	96.6	0.3
" ..	92.3	92.1	-0.2
" ..	91.3	90.9	-0.4
" ..	84.4	83.7	-0.7
" ..	91.6	91.5	-0.1
" ..	84.8	84.7	-0.1
Raw ..	91.7	91.8	0.1
Refined ..	81.7	81.7	0.0
Muscavado ..	88.7	88.7	0.0
Centrifugal ..	96.3	96.5	0.2

The greatest deviation shown in this table is in the case of two sugars, in which the difference is 0.7. This is not due to impurities affecting the saccharometer, as I found by inverting the solution of the sugar in each case. In one of the sugars I found quite a number of soft lumps, which showed that the sugar was very uneven. In the other sugar nothing was found to account for the discrepancy.

The alcoholometers used in these tests are those of Tralles, which are used by the United States Government. These do not differ materially from those of Gay-Lussac. They are more apt to differ from one another on account of defective construction, than they should do from Gay-Lussac's instrument.

Alcoholometers, like other areometers, are very unreliable, unless placed in the hands of those who understand their use. We may, however, use alcoholometers that are not strictly accurate, if only the divisions of the scale bear to one another the proper proportion, and if the displacement of the scale is not too great. This does not apply to the preparation of the standard solution, which should stand correctly at 77.1, which degree of the alcoholometer of Tralles corresponds to specific gravity 0.87.

When we use an alcoholometer that is properly graduated, and we have the standard solution standing at 77.1, we should add 22.9 to the degree of the alcoholometer at 15° C., to obtain the percentage of sugar. If the alcoholometer, through being defective, should not stand at 77.1 for 0.87 specific gravity, but at 76.5, the difference between this latter number and 100, being 23.5, this is the number to be added to the reading of the areometer at 15° C., to obtain the percentage of sugar in the sample under examination.

I have said that cane sugars of all grades, from the highest to the lowest, have given satisfactory results by this process. This assertion is borne out by the examples in table No. 3. I wish, however, to call attention to this, that I do comprise, under the head of sugars, melados or molasses. As to melados, those that have drained out and are comparatively dry may be classed as sugars, but wet melados give very poor results. This may be very easily understood if we remember what was said of the action of water in lowering the alcoholometric degree of a saturated solution, as exemplified in table No. 1. I have little doubt, however, that by experimenting with saturated solutions of methylic hydrated of higher alcoholic degree, we may obtain a solution specially adapted for testing wet melados. It might even be possible, with methylic hydrate of still higher alcoholometric degree, to test molasses within certain limits of density. As an operator can always tell beforehand whether what he has to test is properly a sugar, a wet melado, or a sample of molasses, he could always use the standard solution which was the best adapted to his sample.*

* With melados and molasses, as well as with very low sugars, the direct test seldom gives the true percentage of sugar. It is necessary

From past experience, with saturated solutions of various strengths, I may venture the opinion that methylic alcohol of 87° of the alcoholometer, saturated with sugar, would make a good standard solution for wet melados. For molasses of density about 40° B., methylic alcohol of 90.1° (alcoholometer), saturated with sugar, would not, as far as I can judge without trying it, be far from the proper strength. These are merely hints thrown out to chemists who are interested in testing sugar. Should I not work out these suggestions, I hope that others may find time to study them. The advantages of a process that allows a commercial saccharine product to be tested in about five minutes, are too great to be neglected. The weights of melados or molasses to be taken, would have to be determined, as was done for the weight 19.8 grms., as shown in table No. 2.

(To be continued.)

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 7.

Action of Antimony Trichloride and Tin Tetrachloride upon Naphthalin and other Aromatic Hydrocarbons.—Watson Smith.—In this preliminary communication the author states that he obtains a compound in which the chlorine of antimony trichloride is completely replaced either by three naphthalin residues or by a naphthalin residue and an atom of oxygen. He did not succeed in obtaining this substance in quantity.

Examination of an Aniline Residue.—C. Hell and P. Schoop.—The solid portion of the residue consisted of para-toluylen-diamin. The liquid portion is likewise to a great extent composed of toluylen-diamin, either in a new modification or rendered incapable of crystallising by the presence of isomers.

Addition Products of Acetic Acid with Bromine and Hydrobromic Acid.—C. Hell and O. Mühlhäuser.—The authors have confirmed their results formerly communicated, and determined the boundaries within which the combinations of acetic acid, bromine, and hydrobromic acid are possible.

An Addition Product of Acetic Acid with Bromine and Hydrochloric Acid, and the Power of Acetic Acid to Absorb Bromine and Hydrochloric Acid.—C. Hell and O. Mühlhäuser.—An appendix to the foregoing paper.

Action of Bromine upon Acetic Acid.—C. Hell and O. Mühlhäuser.—The quantities of hydrobromic acid formed increase in geometrical progression up to the formation of a crystalline addition product.

Chloral Hydrate.—A. Naumann.—The author endeavours to prove the resolution of this compound into chloral and water by fractional distillation, and examining the composition of the distillates and the residues.

Specific Heat of Urano-uranic Oxide and the Atomic Weight of Uranium.—Julius Donath.—The average of four determinations of the specific heat of this

to invert, to obtain the correct result, and when we invert a solution that has been imperfectly decolorised, it becomes so red, after inversion, that it is a difficult matter to see through the tube. With chloride of tin and filtration over bone-black, it is possible to improve the colour of the solutions, but even these often fail. The bone-black should be dry and not too great a quantity should be used. On account of these difficulties with dark solutions, experiments to establish the strength of the standard solution and the weight of sugar may have to be made entirely with refined syrups or with low grade refined sugars, to which 5 or 6 per cent additional of water should be added.

compound is 0.07979. The atomic weight of uranium is 120 and the atomic heat 5.96.

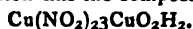
Preparation of Barium from Barium Amalgam.—Julius Donath.—The author criticises the methods proposed by Crookes, Bunsen, and S. Kern, and considers that the metal thus obtained is in all cases contaminated with mercury, pure barium being only obtained electrolytically.

Oxidation of Chinolin by Potassium Permanganate.—S. Hoogewerff and W. A. van Dorp.—The authors obtain by this reaction dicarbopyridic acid, thus confirming the view of a close connection between chinolin and pyridin.

Preparation of Mono- and Di-substituted Organic Malonic Acids.—M. Conrad.—Not susceptible of useful abstraction.

Possibility of enabling Chloro-phyllaceous Plants not known as Saprophytic or Parasitical, to Dispense with the Carbonic Acid of the Air by Means of a Supply of Organic Matter.—M. Schmöger.—Stutzer has answered this question in the affirmative. The author does not consider his experiments decisive.

A New Copper Nitrite.—B. van der Meulen.—The compound in question has the composition—



Quantitative Determination of Cadmium.—F. Beilstein and L. Jawein.—The authors dissolve the precipitated cadmium sulphide (or oxide) in nitric acid, neutralise with potassa, and add potassium cyanide till the precipitate is just re-dissolved. The liquid is then if necessary diluted with water so that 75 c.c. may contain about 0.2 grm. cadmium. The beaker with the solution is then placed in a capsule with cold water, the platinum electrodes are introduced, and the glass is carefully covered to prevent loss by spitting. Three Bunsen elements are required for precipitation. Towards the end of the experiment the glass cover, the electrodes, and the sides of the beaker are rinsed down into the liquid, and the current is allowed to pass for some time longer. The completeness of the precipitation may be ascertained by testing the liquid with sulphuretted hydrogen. The precipitated cadmium is washed first with water, then with alcohol, and dried in a heated platinum capsule.

Action of Alcoholic Solutions of Potassa upon α -Dinitro-chlor-benzol Dissolved in the same kind of Alcohol. Preparation of α -Dinitro-phenylen-ethyl-ether, of α -Dinitro-phenyl-allyl-glycerin, and Phenyl-ether.—C. Willgerodt.—This paper does not admit of useful abstraction.

Action of Basic Bodies upon α -Dinitro-chlor-benzol in a Solution of Carbon Disulphide. Preparation of Dialpha-dinitro-phenyl-sulphide.—C. Willgerodt.—The author obtains yellow, sparingly soluble, pulverulent bodies, which do not melt at 280° to 300°, but generally explode at higher temperatures.

Action of Hydrocyanic and Hydrochloric Acids upon Methyl-acetic Ether.—H. König.—The author obtains oxyadipic acid by a reaction similar to that by which Demarcay prepares oxypropyrotartaric acid.

Contribution to a Knowledge of the Substituted Nitrogen Chlorides.—H. Köhler.—The author supports the view of Wurtz that ethyl-amin-bichloride is truly an ethylated nitrogen chloride.

Aromatic Sulpho-ureas.—B. Rathke.—Both mono- and diphenyl-sulpho-urea dissolve in aqueous potassa and soda (but not in liquid ammonia), and are re-precipitated unchanged by acids, even in carbonic. The same rule holds good with phenyl-xanthogenamid.

Action of Phenylated Oil of Mustard upon Diphenyl-guanidin.—B. Rathke.—By this reaction the author obtains a new base readily soluble in chloroform, but sparingly in benzol and alcohol. The author has been prevented by ill health from completing its examination.

On Biguanide.—B. Rathke.—A very complete description of this new base, $\text{C}_2\text{N}_5\text{H}_7$, which the author considers as formed by the union of equal molecules of guanidin and cyanimide.

Preparation and Analysis of Potassium Ultramarine.—Karl Heumann.—The author obtains this compound as a blue powder by the reaction of an excess of melting potassium chloride or iodide upon silver ultramarine.

The Thermo-mechanical Relation between the Boiling-point and the Melting-point of the Solid Elements.—H. F. Wiebe.—The author undertakes to show that for a series of bodies that quantity of heat which is requisite at the constant pressure of one atmosphere to raise equal volumes of the same from the melting-point to the boiling-point bears the same constant proportion to their inverse absolute coefficients of expansion.

Remarks on O. Dœbner's Paper on "Malachite Green."—E. and O. Fischer.—This paper relates, first, to the personal question of the priority of discovery, and, secondly, to the constitution of colouring-matter.

Colouring-Matters of the Rosanilin Group.—E. and O. Fischer.—An account of benzoyl-green, of its reduction, and of its para- and meta-nitro compounds.

On Para-cresol.—E. Baumann and L. Brieger.—Paracresol is the chief constituent of the mixture of phenols obtained during the putrefaction of albumen, and regarded hitherto exclusively as phenol. It is also the chief constituent of the volatile phenols obtained on distilling the urine of horses with hydrochloric acid.

Brom-phenyl-mercapturic Acid.—E. Baumann and C. Preusse.—The authors describe the results of an experiment where a dog received daily doses of brom-benzol. The urine was found to contain sulpho-brom-phenolic acid in abundance. They then describe the decomposition of brom-phenyl-mercapturic acid by boiling with alkalis and by acids.

A Series of Homologous Tertiary Diamines observed in the Manufacture of Methyl-anilin.—O. Dœbner.—The author describes the first member of this series, $\text{C}_{17}\text{H}_{22}\text{N}_2$. A base, $\text{C}_{19}\text{H}_{26}\text{N}_2$, examined by A. W. Hofmann and Martius belongs to the same series.

Benzyl-methyl-glycolic Acid.—S. Gabriel and A. Michael.—The authors have been engaged with the attempt to prepare benzyl-methyl-glycolic acid by the action of hydrocyanic and hydrochloric acids upon phenyl-aceton, and then to reduce it to benzyl-methyl-acetic acid.

Oxytoluylic Acids obtained from the three Isomeric Cresols by means of Carbon Tetrachloride, and their Oxidation to Oxyphthalic Acids.—C. Schall.—This paper does not admit of useful abstraction.

Textile Colorist. Vol. i., No. 3.

This issue contains a variety of useful matter. The history of aniline by P. Prunier, described as being copyright, contains very little indeed that has not gone the round of chemical manuals. Articles borrowed from other journals are acknowledged in a somewhat unsatisfactory manner. Thus, certain instructions for testing the dyes of coloured fabrics are said to be from the *Gazette*, and a paper on aniline-black is ascribed to the *Bulletin*. Such references seem to us too vague. In the last-mentioned paper we read:—"A queer shade is thus obtained, which, after washing and scouring, becomes a pure black." We should suppose that a green shade is meant. Further on a patent process is described as involving the use of a solution of "ferruginous chlorine." Attention to correctness in technical terms will greatly increase the value of the paper.

MISCELLANEOUS.

The British Association.—At the concluding meeting of the British Association at Sheffield, Captain Galton, having read the report of the General Committee, said that the numbers attending the meetings of the Association were, old life members 184, new life members 16, old annual members 239, new annual members 74, associates 529, ladies 349, foreign members 13, total 1404, and the amount in money was £1425. The grants, amounting to £960 were appropriated as follows:—

A.—MATHEMATICS AND PHYSICS.

	£	s.	d.
Dr. Lodge—New Form of High Insulation Key	10	0	0
Prof. Adams—Standard of White Light ..	20	0	0
Prof. Everett—Underground Temperature ..	10	0	0
Dr. Joule—Determination of the Mechanical Equivalent of Heat ..	50	0	0
Sir. W. Thomson—Elasticity of Wires ..	50	0	0
Mr. Glashier—Luminous Meteors ..	30	0	0
Mr. G. H. Darwin—Lunar Disturbance of Gravity ..	30	0	0
Prof. Sylvester—Fundamental Invariants ..	50	0	0
Mr. J. Perry—Laws of Water Friction ..	20	0	0
Prof. W. E. Ayerton—Specific Inductive Capacity of Sprengel Vacuum ..	20	0	0
Rev. Prof. Haughton—Completion of Tables of Sun-heat Coefficients ..	50	0	0
Prof. G. Forbes—Instruments for Detection of Fire-damp in Mines ..	10	0	0
Mr. J. M. Thomson—Inductive Capacity of Crystals and Paraffins ..	25	0	0

B.—CHEMISTRY.

Prof. Dewar—Spectrum Analysis ..	10	0	0
Dr. Wallace—Development of Light from Coal-gas ..	10	0	0

C.—GEOLOGY.

Prof. P. M. Duncan—Report on Carboniferous Polyzoa ..	10	0	0
Prof. A. L. Adam—Caves of South Ireland ..	10	0	0
Prof. Seeley—Viviparous Nature of Ichthyosaurus ..	10	0	0
Mr. John Evans—Kent's Cavern Exploration ..	50	0	0
Mr. John Evans—Geological Record ..	100	0	0
Prof. W. C. Williamson—Miocene Flora of the Basalt of North Ireland ..	15	0	0
Prof. Hull—Underground Waters of Permian Formations ..	5	0	0

D.—BIOLOGY.

Dr. Pye-Smith—Elimination of Nitrogen by Bodily Exercise ..	50	0	0
General M. Lane-Fox—Anthropological Notes	20	0	0
Mr. Stainton—Record of Zoological Literature	100	0	0
Dr. M. Foster—Table at Zoological Station at Naples ..	75	0	0
Dr. A. Gamgee—Investigation of the Geology and Zoology of Mexico ..	50	0	0
Sir J. Lubbock—Excavations at Port Stewart	15	0	0

E.—STATISTICS AND ECONOMIC SCIENCE.

Dr. Farr—Anthropometry ..	50	0	0
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G.—MECHANICS.

Mr. Bramwell—Patent Laws ..	5	0	0
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The meeting in 1880, will be held at Swansea commencing on August 25, under Presidency of Prof. Ramsay, LL.D., F.R.I. In 1881 the Association will revisit York, where the first meeting, in 1831, was held.

University of London.—The following is a list of the candidates who have passed the recent Examinations for Honours, First B.Sc., and Preliminary Scientific (M.B.) Examinations:—

(First B.A. and First B.Sc. conjointly.)

Mathematics—First Class: S. Loney, First B.A.,

Sidney Sussex College, Cambridge. Second Class: Upen-
dra Kriahna Dutt, First B.Sc. and Prel. Sci., University
College. Third Class: T. G. Creak, First B.A., Owens
College.

(First B.Sc. and Preliminary M.B. conjointly.)

Chemistry—First Class: A. P. Luff, Prel. Sci., St. Mary's Hospital; J. S. W. Chitty, First B.Sc., Magdalen College, Oxford; W. A. Gostling, Prel. Sci., University College. Second Class: W. L. Goodwin, First B.Sc., University of Edinburgh; Frances H. Prideaux, Prel. Sci., London School of Medicine for Women; Emily Tomlinson, Prel. Sci., Girton College, Cambridge. Third Class: A. K. A. Spiegel, First B.Sc., Owens College; W. E. Wynter, Prel. Sci., St. Bartholomew's and Middlesex Hospitals; H. Duncan, Prel. Sci., University College; J. Stevenson, Prel. Sci., Owens College; P. F. Frankland, First B.Sc., Royal School of Mines and private study; A. J. Turner, Prel. Sci., University College.

Experimental Physics—First Class: J. Ryan, First B.Sc. (disqualified by age for the Arnott Medal), Cambridge (unattached). Second Class: H. Duncan, Prel. Sci., University College; W. A. Slater, Prel. Sci., Guy's Hospital; E. P. Cockey, Prel. Sci., Epsom College; S. Young, First B.Sc., Owens College; W. L. Goodwin, First B.Sc., University of Edinburgh; A. J. Turner, Prel. Sci., University College; F. Womack, Prel. Sci., St. Bartholomew's Hospital; R. B. Lee, First B.Sc., private study. Third Class: W. E. Wynter, Prel. Sci., St. Bartholomew's and Middlesex Hospitals; P. P. Whitcombe, Prel. Sci., Epsom College and St. Mary's Hospital; T. R. Dallmeyer, First B.Sc., University College and private study; S. F. Harmer, First B.Sc., University College; H. Settle, Prel. Sci., St. Bartholomew's Hospital; C. S. Spong, Prel. Sci., Epsom College and Guy's Hospital.

Botany—First Class: A. M. Vann, Prel. Sci., King's College; G. W. Hill, First B.Sc., King's College and St. George's Hospital. Second Class: R. F. Fox, Prel. Sci., private study and London Hospital; Catherine Alice Raison, First B.Sc., private study; T. S. Short, Prel. Sci., King's College.

Zoology—First Class: B. F. Halford, Prel. Sci. (Exhibition), University College; A. J. Turner, Prel. Sci., University College. Second Class: T. W. Shore, Prel. Sci., Hartley Institute and Royal School of Mines. Third Class: S. S. Merrifield, Prel. Sci., King's College; A. E. Taylor, Prel. Sci., University College.

(First M.B. Examination.)

Organic Chemistry—First Class: E. L. Adeney (Exhibition and Gold Medal), Guy's Hospital; *W. Lane, Guy's Hospital; C. P. Lukis, St. Bartholomew's Hospital. Second Class: J. J. Udale, Guy's Hospital. Third Class: R. Prothero, Liverpool School of Medicine and Guy's Hospital; R. Sisley, St. George's Hospital.

Materia Medica and Pharmaceutical Chemistry—First Class: A. Barron (Exhibition and Gold Medal), Owens College and Liverpool Royal Infirmary; †A. Daniell, Universities of Edinburgh and Paris; W. T. Maddison, King's College. Second Class: A. G. Dawson, Owens College; J. M. Rogers, Middlesex Hospital; T. G. Stenham, London Hospital; G. C. R. Bull, St. Mary's Hospital; W. E. Fielden, Guy's Hospital.

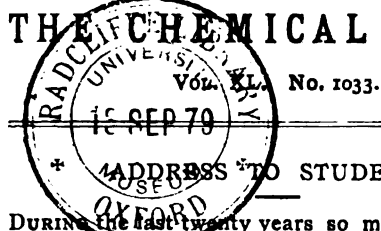
* Obtained the number of marks qualifying for a Medal.

† Obtained the number of marks qualifying for the Exhibition.

NOTICE.

The STUDENTS' NUMBER of the CHEMICAL NEWS will be published on Friday, September 12th. Gentlemen holding official positions in the Universities, Colleges, Medical Schools, &c., of the United Kingdom, where Chemistry and Physical Science form a part of the education, who have not yet forwarded the necessary information to our Office for publication in that Number, will confer a favour by sending it with the least possible delay.

THE CHEMICAL NEWS.



No. 1033.

DURING the last twenty years so many "Addresses to Students" have appeared in the CHEMICAL NEWS as prefaces to the annual "Students' Number" that it has become difficult—nay, almost impossible—to say anything new. We have done our best—but, alas! unsuccessfully—to exorcise that evil spirit "Cram;" we have steadily advocated the necessity of wedding practice to theory by indissoluble bonds; we have warned the warm-blooded young student against theoretical Will-o'-the-wisps and formulistic Jack-o'-Lanterns; and we have tried to prove to him the fallacy of trying to follow too many paths at once, by showing that all truly great philosophers, whether chemists or physicists, were men of a single and steadily followed aim, until we have worn every subject to shreds. We have, therefore, resolved to be personally silent on the present occasion, and intend to confine ourselves to picking out a few passages from a speech made by that Ithuriel of the "Cram Demon," Prof. Huxley, to the London University College schoolboys at the annual distribution of prizes on July 31st. Although delivered to boys still at school, it contains a notable amount of good advice that may be taken to heart by students of all ages.

The beginning of the Professor's address was a telling sketch of the kind of boys who win prizes, a sketch which will equally apply to men who are successful in carrying off the more or less solid honours of the University. Quickness in learning, readiness and accuracy in reproducing, industry, and, above all, endurance, are the essential characteristics of the prize winner *par excellence*. We may, however, venture to suggest that Prof. Huxley describes the typical prize winner as he would like to see him rather than as he is. As an instance of this it may be stated that not so very long ago a certain student, who had never handled a dissecting knife, carried off the B.A. prize, as well as honours in Animal Physiology, by dint of an excellent memory. Prof. Huxley justly places the power of bearing both mental and physical fatigue in the very front of the qualifications of a successful student, and reminded his hearers that this power is not necessarily the concomitant of either great mental or physical strength. To a lawyer, physician, or merchant, the power of being able to work for fifteen or sixteen hours at a stretch without being knocked up may make all the difference between success and failure.

College life once past, Prof. Huxley asks from the would-be successful competitor as much industry as he can bring to bear on his particular subject, a broad deep chest, and a stomach of whose existence he knows nothing. Competitive examination is only a partial test of what the prize winners will be in after life. The student just referred to is now a Government clerk; and a contemporary of his who in the same year carried off prizes and honours in chemistry without ever having cleaned a test-tube in his life, is now one of our leading musical com-

posers and critics. A more practical and instructive commentary on the evils of "cram" could hardly be found than is afforded us by these two cases. The real gist of the matter is that those who fail at college are more than frequently successful competitors in the great struggle for a place in the world's lists. "Those," said Prof. Huxley, "who have won prizes have made a good beginning; those who have not, may yet make that good ending which is better than a good beginning. Professor Circumstance is the severe pedagogue who conducts the long competitive examination of practical life."

Let those, therefore, who not only win no prizes, but who fail even to pass, take heart of grace. The winners of rewards and places unquestionably give promise of doing great things in the future. But how many of them have become distinguished members of society? Whilst among the undistinguished crowd there may be some simple student with whose name the world may one day ring. The unsuccessful ones, then, who go down before their more brilliant compeers should remember that in the battle of life the steady-burning farthing rushlight is of more real value than the dazzling meteor which flashes across half the firmament. Win prizes and honours if you can. If you have the other qualities necessary for success in this world, such distinctions will only act on you as a mental tonic, a sharp but necessary spur to goad you on to reach the goal; but if, in spite of all your industry and perseverance, you find yourself at the bottom of the list, comfort yourself with the feeling that as long as you possess industry, honesty, and courage, the greater and more enduring prizes of the world are still open to you.

Twenty years ago the late J. G. Edgar wrote a charming boy's book, entitled the "Boyhood of Great Men." For the encouragement of those who have been unsuccessful in carrying off prizes at school or college, will no one write a companion to this, to be called the "Manhood of Great Boys?"

In a leader on the subject of competitive examinations and the cramming system generally, the *Times* makes the following apposite and incisive remarks:—

"Several circumstances have combined of late to direct public attention to the subject of examinations. Cambridge, as we saw a short time ago, is forced seriously to consider the system of entrance scholarships; and a correspondent has recently invited attention to the mischievous system of entrance scholarships established in public schools. Even if examinations were a good in themselves, it would still be possible to have too much of a good thing, and very many competent judges are already beginning to dread an educational surfeit in this respect. But, to our thinking, examinations, so far from being a good in themselves, can hardly be placed higher than the category of necessary evils. They are a burden to the examiner and to the teacher, exactly in proportion as each is efficient and conscientious, and they are very far from being an unalloyed benefit even to the examinee. But they are not the less in many cases necessary; they are at least a rough test of merit, capacity, and attainment, and therefore, where these have to be tested for any specific purpose, it is hardly possible to dispense with examination altogether. But to admit this much is very far from saying that examinations should be made, what they are fast becoming, the be-all and end-all of educational processes. From the tender age of ten or eleven to that of manhood and upwards nearly every boy of promising parts in this country lives with the constant fear of examinations before his eyes. If he is more than usually suc-

cessful, at the end of the process he is likely to enjoy for the next few years, or, indeed, as long as he chooses, the distinguished privilege of examining his juniors in their turn. Thus the examination fever spreads far and wide. It spares neither age nor sex; for women, with singular perversity, have claimed as a privilege what boys and men alike regard with aversion. It has long ago pervaded education, and its contagion is now beginning to infect the whole range of modern letters. Literature, ancient and modern, is regarded as so much material for examination to be reproduced in the form best calculated to win marks in a competition. History is cut up into "periods" and "epochs," and then reduced into summaries, so that whoso runs may read or teach, examine or be examined. The old Universities, which once could boast of a learned press and still occasionally publish works not unworthy of English scholarship, devote their chief literary energies to the publication of manuals required in the various examinations they have undertaken to conduct. The work is excellently done, no doubt, though it is hardly of a kind which befits the dignity of an Academical press. But the examination spirit is rampant, and the Universities are forced to yield to it. The pity of it is that they take to the task so kindly and seem so entirely contented with it.

"The evil of all this is unquestionably great and growing. We need not dwell on the disastrous effects, well known to schoolmasters and college tutors, of the premature forcing for the purpose of competitive examination to which so many boys of tender years are now submitted in public schools. It is rather to the general effects of the modern tendency to make examination the sole test and crown of all processes of education that we wish to draw attention. This tendency entirely distorts every rational view of what education is and should be. It makes of the pupil a mere racer, and one who contends for heavy pecuniary stakes. It makes of the teacher a trainer whose whole prosperity depends, not on his power of imparting sound knowledge and drawing out the natural capacities of the mind, but on his skill in preparing his pupils for a particular competition. It makes of the examiner a judge, not of mental capacity and sound information generally, but of those qualities alone which are readily estimated in marks. In addition to all this the system tends inevitably to force teaching and examining alike into a narrow and mechanical groove. Even if a particular teacher has a special taste and regard for some subject out of the ordinary range of the examination for which he is preparing his pupils, he dare not lead them in the direction in which he would probably do them most good, for fear they should fail to get credit for their work in the coming ordeal, on which their whole success in life may depend. He is forced to scan with anxious scrutiny the line that the examination has previously taken, in the confidence, very rarely misplaced, that it will take the same line again. For the examiner knows that he, too, must not go beyond certain well-understood limits. If he does, he will be regarded as crotchety, unfair, and pedantic. Every experienced teacher knows to his cost how the attempt to lead his pupils towards some collateral line of study not directly recognised in an examination is frustrated at once by their refusal to take any interest in subjects that will not "pay." Every examiner knows that the insertion in his papers of a question lying somewhat out of the recognised range and groove is simply so much waste of time and labour. Hence, under the influence of examinations, the treatment of every subject, great and small, is divided, by an impassable barrier, into the dark and boundless range of the neglected and the unknown, which the teacher must leave unnoticed and the examiner dares not explore, and the narrow field, brilliantly illuminated and minutely surveyed, which the pupil is taught to regard as alone worthy of notice. Even thus we have not exhausted the evil effects of thus substituting examination for education. The whole system gives the successful competitors an exaggerated sense of

the importance of the victories they have won. It unduly stimulates their earlier efforts, while it paralyses their later and more mature energies. A high wrangler or a first-class man thinks that the battle of life is won. He has learnt all that he can learn, has done all that he needs to do. He despises knowledge which lies outside the examination range as musty, pedantic, and unprofitable. He thinks meanly of men who have not been examined so often or so successfully as himself. He owes all that he is, and has, and knows to having been examined: he believes in the process, and he aims at nothing higher than being in his turn an examiner himself. Then, indeed, his fate is sealed: he might have been a student, a scholar, or a philosopher if he had not been taught to look at all knowledge through the distorting medium of examinations; he might even have succeeded in life, in spite of early obstacles and mischievous training, if he had not been led to believe that that success was already won at its outset. But he becomes a mere subordinate wheel in a vast and exacting machine; his existence is passed in a weary and monotonous round of setting papers and looking them over, of assigning marks and adding up their total, of comparing results with his colleagues and striving to gauge human nature by impossible measurements and fallacious standards, until at last he comes to believe that there is no better fate in store for human beings than to become just what he is himself.

"The picture we have drawn is highly coloured, perhaps, but it will the better serve to illustrate the inevitable consequences of making examination supreme, instead of keeping it in proper subordination to the higher purposes of education. It used to be said that in some districts of England a man could be supported by charity from his cradle to his grave. We have changed all that, and in the place of indiscriminate charity we have established the supremacy of examinations at every turn in life. Is the result so entirely satisfactory that we can regard it as final and irrevocable? Is it not possible that, as so often happens, we have confounded ends with means and made a successful examination the paramount purpose, instead of merely the indispensable test, of educational training? What, in fact, is the actual result? Education is doubtless improved in its lower ranges, and the public service of the country has been purged of mere favour and nepotism. These are valuable and indisputable gains; but they would be purchased dearly by the sacrifice of high aims in academical culture and the absorption of some of the best capacity of the country in a kind of work which extinguishes original study and paralyses all intellectual independence. If we persist in absorbing some of the best capacity of Oxford and Cambridge in the perpetual conduct of trivial and elementary examinations, can there be any reasonable chance of those Universities producing results proportionate to their fame and endowments in the real work of learning?"

Such sentiments coming from the leading journal will be read with gratification by all true friends of education, who will rejoice to see the *Times* joining in the battle against the cram system, a battle which the *CHEMICAL NEWS* has fought unflinchingly from its first number.

UNIVERSITIES AND COLLEGES.

UNIVERSITY OF LONDON.

CANDIDATES for any Degree granted by this University are required to have passed the Matriculation Examination, to which no candidate is admitted unless he has produced a certificate showing that he has completed his sixteenth year.

The Fee for this examination is £2.

The Examination will be held on Monday, January 12th, 1880. It is conducted by means of Printed Papers; but the Examiners are not precluded from putting, for the purpose of ascertaining the competence of the Candidates to pass *viva voce* questions to any Candidate in the subjects in which they are appointed to examine.

Candidates are not approved by the Examiners unless they have shown a competent knowledge in each of the following subjects:—Latin. Any two of the following Languages:—Greek, French, German, and either Sanskrit or Arabic. Arithmetic and Algebra. Geometry. The English Language and English History. Natural Philosophy. Chemistry.

The Papers in Latin and Greek will contain passages to be translated into English, with questions in Grammar and History and Geography arising out of the subjects of the books selected. Short and easy passages will also be set for translation from other books not so selected. A separate paper will be set containing questions in Latin Grammar, with simple and easy sentences of English to be translated into Latin.

Candidates may substitute German for Greek.

The papers in French and German will contain passages for translation into English, and questions in Grammar, limited to the Accidence. The paper in Sanskrit or Arabic will contain passages for translation into English, and questions in Grammar.

The examination in the English Language and English History includes Orthography; Writing from Dictation; the Grammatical Structure of the Language.

History of England to the end of the Seventeenth Century.

That in Mathematics includes the ordinary Rules of Arithmetic; Vulgar and Decimal Fractions; Extraction of the Square Root.

Addition, Subtraction, Multiplication, and Division of Algebraical Quantities; Proportion; Arithmetical and Geometrical Progression; Simple Equations.

The First Four Books of Euclid, or the subjects thereof.

The Questions in Natural Philosophy are of a strictly elementary character; they include Mechanics, Hydrostatics, Hydraulics, Pneumatics, Optics, and Heat.

The Examination in Chemistry is—Chemistry of the Non-metallic Elements; including their compounds—their chief physical and chemical characters—their preparation—and their characteristic tests.

A Pass Certificate, signed by the Registrar, will be delivered to each Candidate who applies for it, after the Report of the Examiners has been approved by the Senate.

If in the opinion of the Examiners any Candidates in the Honours Division of not more than Twenty years of age at the commencement of the Examination possess sufficient merit, the first among such Candidates will receive an Exhibition of thirty pounds per annum for the next two years; the second among such Candidates will receive an Exhibition of twenty pounds per annum for the next two years; and the third will receive an Exhibition of fifteen pounds per annum for the next two years; such exhibitions are payable in quarterly instalments, provided that on receiving each instalment the Exhibitioner declares his intention of presenting himself either at the two Examinations for B.A., or at the two Examinations for B.Sc., or at the First LL.B. Examination, or at the Preliminary Scientific and First M.B. Examinations, within three academical years* from the time of his passing the Matriculation Examination.

* By the term "Academical Year" is ordinarily meant the period intervening between any Examination and an Examination of a higher grade in the following year; which period may be either more or less than a Calendar year. Thus the interval between the First Examinations in Arts, Science, and Medicine, and the Second Examinations of the next year in those Faculties respectively, is about sixteen months, whilst the interval between the Second B.A. Examination and the M.A. Examination of the next year, or between the Second B.Sc. Examination and the D.Sc. Examination of the next year, is less than eight months. Nevertheless, each of these intervals is counted as an "Academical Year."

Under the same circumstances, the fourth among such Candidates will receive a prize to the value of ten pounds in books, philosophical instruments, or money; and the fifth and sixth will each receive a prize to the value of five pounds in books, philosophical instruments, or money.

Any Candidate who may obtain a place in the Honours Division at the Matriculation Examination in January is admissible to the First B.A. or to the First B.Sc. Examination in the following July. But such Candidate will not be admissible to the Second B.A. or to the Second B.Sc. Examination in the ensuing year, unless he has attained the age of eighteen years.

FIRST B.Sc. EXAMINATION.

The First B.Sc. Examination will be held in July, 1880.

No Candidate (with the exception of such as have obtained Honours at the Matriculation Examination in the preceding January) is admitted to this Examination within one academical year of the time of his passing the Matriculation Examination.

The Fee for this Examination is £5.

The Examination embraces the following subjects:—Pure and Mixed Mathematics, Inorganic Chemistry, Experimental Physics, and General Biology.

Examination for Honours.

Any Candidate who has passed the First B.Sc. Examination in all its subjects may be examined at the Honours Examination next following the First B.Sc. Examination at which he has passed for Honours in (1) Mathematics, (2) Experimental Physics, (3) Chemistry, (4) Botany, and (5) Zoology; unless he has previously obtained the Exhibition in Pure and Mixed Mathematics at the First B.A. Examination, in which case he will not be admissible to the Examination for Honours in that subject; or unless he has previously obtained the Exhibition at the Preliminary Scientific (M.B.) Examination in either of the subjects which are common to it with the first B.Sc. Examination, in which case he will not be admissible to the Examination for Honours in that subject.

Candidates for Honours in Chemistry will be examined in Inorganic Chemistry, treated more fully than in the Pass Examination. In addition, they will be examined practically in Simple Qualitative and Quantitative Analysis. This Examination, which will consist of six hours' examination by printed papers and of six hours' practical work, will take place on Thursday and Friday in the same week with the Examination for Honours in Mathematics, commencing on each day at 10 a.m.

In the Examination for Honours, the Candidate, not being more than 22 years of age at the commencement of the Pass Examination, who most distinguishes himself in Chemistry or Experimental Physics, will receive an Exhibition of £40 per annum for the next two years.

SECOND B.Sc. EXAMINATION.

The Second B.Sc. Examination will be held in October, 1880.

Candidates for this Examination are required to have passed the First B.Sc. Examination at least one academical year previously.

The Fee for this Examination is £5.

The regulations are framed with the view of allowing the candidate to bring up *any three* of the following nine subjects:—

1. Pure Mathematics.
2. Mixed Mathematics.
3. Experimental Physics.
4. Chemistry.
5. Botany, including Vegetable Physiology.
6. Zoology.
7. Animal Physiology.
8. Physical Geography and Geology.
9. Logic and Psychology.

It is understood the amount of proficiency expected in each of the three subjects chosen will be that which the candidate might attain by the steady devotion to it of about one-third of the sessional work of a diligent student.

Examination for Honours.

Any Candidate who has passed the Second B.Sc. Examination, and has not previously passed the Second B.A. Examination, may be examined at the Honours Examination next following the Second B.Sc. Examination at which he has passed, for Honours in (1) Mathematics, (2) Logic and Psychology, (3) Experimental Physics, (4) Chemistry, (5) Botany, (6) Zoology, (7) Physiology, (8) Physical Geography and Geology; provided that he shall have gone through the Pass Examination in the corresponding subject or subjects immediately before. And any Bachelor of Arts who has passed the Second B.Sc. Examination may under the same conditions be examined for Honours in one or more of the above mentioned subjects, unless he have previously obtained a Scholarship at the Second B.A. Examination in either of the first two of those subjects, in which case he shall not be admissible to the Examination for Honours in that subject.

The examination for Honours in Chemistry will take place on Monday and Tuesday in the second week after the conclusion of the Pass Examination; on Monday by printed papers (chiefly on Organic Chemistry), and on Tuesday by practical exercises in Simple Qualitative and Quantitative Analysis.

The candidate, being not more than 23 years of age, who most distinguishes himself in Chemistry, will receive £50 per annum for the next two years, with the style of University Scholar.

DOCTOR OF SCIENCE.

The examination for the Degree of Doctor of Science takes place annually within the first twenty-one days of June, and the examination in each branch occupies four days.

No candidate is admitted to the examination for the Degree of D.Sc. until after the expiration of two Academic Years from the time of his obtaining the Degree of B.Sc. in this University, unless he shall have passed the Second B.Sc. Examination in the First Division at least two Academic years subsequently to having passed the first B.Sc. Examination, in which case he shall be admitted to the examination for the Degree of Doctor in Science at the expiration of one Academic Year from the time of obtaining his B.Sc. Degree.

The Fee for this Examination is £10.

Every candidate for the degree of D.Sc. is examined in some one or more of the various branches of Physical, Biological, or Mental Science, to be selected by himself; and no candidate is approved by the examiners unless he has shown a thorough practical knowledge of the principal subject and a general acquaintance with the subsidiary subject or subjects, specified as belonging to the branch so selected. He is expected to be so fully conversant with the principal subject he may select as to be able to go through any examination test (whether theoretical or practical) of his acquirements in it that can be fairly applied. Candidates, when giving notice, must specify the branch or branches in which they desire to be examined.

BRANCH IV. OF PHYSICAL SCIENCE. INORGANIC CHEMISTRY.

Principal Subject—Inorganic Chemistry.
Subsidiary Subject—Either Organic Chemistry; or Mineralogy, Crystallography, and Chemical Technology in its relations to Inorganic Chemistry.

BRANCH V., ORGANIC CHEMISTRY.

Principal Subject—Organic Chemistry.

Subsidiary Subjects—Either Inorganic Chemistry; or Chemical Technology in its relations to Organic Chemistry, and the Chemistry of Animal and Vegetable Life.

PRELIMINARY SCIENTIFIC (M.B.) EXAMINATION.

This examination will be held in July, 1880.

No Candidate is admitted to this examination until he has completed his seventeenth year, and has either passed the Matriculation Examination* or taken a Degree in Arts in one of the Universities of Sydney, Melbourne, Calcutta, or Madras (provided that Latin was one of the subjects in which he passed). The fee for this examination is five pounds.

Candidates for the degree of M.B. are strongly recommended by the Senate to pass the Preliminary Scientific Examination before commencing their regular medical studies; and to devote a preliminary year to preparation for it according to the following programme:—Winter Session: Experimental Physics; Chemistry (especially Inorganic); Zoology. Summer Session: Practical Chemistry (Inorganic); Botany.

Any candidate who has passed the Preliminary Scientific (M.B.) Examination, may be examined at the Honours Examination next following the Preliminary Scientific Examination at which he has passed, unless he has previously obtained an Exhibition in any one of the subjects at the First B.Sc. Examination, in which case he is not admissible to the Examination for Honours in that subject.

Candidates for Honours in Chemistry are examined in Inorganic Chemistry, treated more fully than in the Pass Examination. In addition they are examined practically in Simple Qualitative and Quantitative Analysis.

EXAMINATION IN SUBJECTS RELATING TO PUBLIC HEALTH.

A Special Examination will be held in December in subjects relating to public health.

No candidate is admitted to this Examination unless he has passed the Second Examination for the Degree of Bachelor of Medicine in this University at least one year previously; nor unless he shall have given notice of his intention to the Registrar at least two calendar months before the commencement of the Examination.

The Fee for this Examination is £5.

Candidates are examined in the following subjects:—

Chemistry and Microscopy, in relation to the examination of Air, Water, and Food.

Meteorology and Geology, as far as they bear on the duties of Health Officers, viz.:—General knowledge of Meteorological Conditions; Reading and Correction of Instruments. General knowledge of Soils; their Conformation and Chemical Composition.

Vital Statistics, in reference to the methods employed for determining the Health of a Community; Birth-rate; Death-rate; Disease-rate; Duration and of Expectancy of Life. Present amount of Mortality, and its causes, in different Communities.

Hygiene.—General principles of Hygiene. Special topics:—Soil. Construction of Dwellings. Conservancy of Cities. Unhealthy Trades. Supply of Food to Cities, and Examination of Food. Disposal of Sewage. Water-supply.

Medicine, in reference to the origin, spread, and method of prevention of Diseases generally, but especially those of the Epidemic class.

Sanitary Engineering, as far as regards the arrangements connected with Water-supply, Sewerage, and Ventilation. A knowledge of the reading of Plans, Sections, Scales, &c.

* Candidates who pass in all the subjects of the Preliminary Scientific (M.B.) Examination, and also pass at the same time in the Pure Mathematics of the First B.Sc. Examination, or who have previously passed the First B.A. Examination, are considered as having passed the First B.Sc. Examination. The attention of such candidate directed to the fact that, under the new regulations for the B.Sc. Degree, this degree may be obtained by passing at the Second B.Sc. Examination in the three Biological subjects only.

Sanitary Law, as far as it relates to the duties of Officer of Health. A knowledge of the powers given under the various Sanitary Acts, as defined in the Instructions issued by the Local Government Board, and of the methods of procedure in special cases.

The Examination, which is both written and practical, extends over four days.

GILCHRIST SCHOLARSHIPS.

1. A Scholarship of the value of Fifty Pounds per annum, and tenable for three years, is annually awarded to the highest among those candidates at the June Matriculation Examination who have been approved by the Principal of University Hall as fit to be received into that Institution with a view to the prosecution of their studies in University College for Graduation in one of the four Faculties of the University of London; provided that such Candidate pass either in the Honours List or in the First Division.—Particulars may be obtained on application to the Principal of University Hall, Gordon Square, W.C.

2. A similar Scholarship is annually awarded to the Candidate from the Royal Medical College, Epsom, who at the June Matriculation Examination stands highest among the Candidates approved by the Head Master of that Institution, and who passes either in the Honours List or in the First Division; on condition of his prosecuting his studies during the tenure of his Scholarship with a view to Graduation in one of the four Faculties of the University of London.—Particulars may be obtained on application to the Secretary of the Royal Medical College, 37, Soho Square, W.

3. A similar amount is annually offered to Candidates intending to pursue, at Owens College, Manchester, their studies for Graduation in one of the four Faculties of the University of London; a single Scholarship of Fifty Pounds per annum for three years being awarded to the highest of those Candidates at the June Matriculation Examination who shall have been previously approved by the Principal of Owens College, provided that he pass in the Honours Division; or, in case no Candidate should so pass, two Scholarships, each of Twenty-five Pounds per annum, being awarded to the two Candidates as aforesaid who shall stand highest in the First Division.—Particulars may be obtained on application to the Principal of Owens College, Manchester.

Particulars of the Colonial and Indian Scholarships may be obtained on application to the Secretary of the Gilchrist Educational Trust, University of London, W.

SCIENCE AND ART DEPARTMENT OF THE COMMITTEE OF COUNCIL ON EDUCATION, SOUTH KENSINGTON.

A sum of money is voted annually by Parliament for scientific instruction in the United Kingdom. The object of the grant is to promote instruction in Science, especially among the industrial classes, by affording a limited and partial aid or stimulus towards the founding and maintenance of Science schools and classes.

In order to place a science school or class in connection with the Science and Art Department, an approved committee, consisting of at least five well known and responsible persons, must be formed.

The following are the sciences towards instruction in which aid is given by the Science and Art Department:—Practical, Plane, and Solid Geometry; Machine Construction and Drawing; Building Construction; Naval Architecture and Drawing; Pure Mathematics; Theoretical Mechanics; Applied Mechanics; Acoustics, Light, and Heat; Magnetism and Electricity; Inorganic Chemistry; Organic Chemistry; Geology; Mineralogy; Animal Physiology; Elementary Botany; Biology, including Animal and Vegetable Morphology and Physiology; Principles of Mining; Metallurgy; Navigation; Nautical Astronomy; Steam; Physiography; Principles of Agriculture.

The aid is given in the form of—1. Public examinations, in which Queen's Prizes are awarded, held at all places complying with certain conditions; 2. Payments on results as tested by these examinations; 3. Scholarships and Exhibitions; 4. Building grants; 5. Grants towards the purchase of fittings, apparatus, &c.; 6. Supplementary grants in certain subjects, and special aid to teachers and students.

The examinations are held about the month of May under the superintendence of the local committees and local officers. The examination papers are prepared by the professional examiners in London. An evening is set apart for one or more subjects, so that the examination in each subject is simultaneous over the whole kingdom.

A packet of examination papers is sent to each local examination secretary, who opens it in the presence of the committee and candidates. The committee is held responsible that no unfair means of any description are used in working the papers, and that the rules of the Department are strictly complied with.

The examinations are of two kinds, but held together, viz.—

a. The Class examinations, of which there are two grades or stages; the first stage or elementary examination, and the second stage or advanced examination. The successful candidates in both stages are divided into 1st and 2nd class.

b. The Honours examination of a highly advanced character. In this there are also two classes.

Any person however taught may sit at any one of these examinations.

The Queen's prizes, consisting of books or instruments, are also given to all candidates who are successful in obtaining a first class in either stage of the class examinations.

The payments on results are made only on account of the instruction of students of the industrial classes, or on account of the instruction of their children. They are—£2 for a first class, and £1 for a second class, in each stage of the class examinations, and £2 and £4 for a second or first class respectively in honours. Special extra payments are made in attendance in organised Science Schools. Special payments are also made for Chemistry.

There are also two forms of scholarship in connection with elementary schools.

a. In the Elementary School Scholarship £5 are granted to the managers of any elementary school for the support of a deserving pupil selected by competition, if they undertake to support him for a year and subscribe £5 for that purpose. The payment of £5 by the Science and Art Department is conditional on the scholar passing in some branch of science at the next May examination.

b. In the Science and Art Scholarship, which is of a more advanced character, a similar contribution of £5 is required on the part of the locality, and a grant of £10 is made by the Department towards the maintenance, for one year, of the most deserving pupil or pupils in elementary schools who have passed certain examinations in science and in drawing.

In both these cases the scholar must be from twelve to sixteen years of age, and one scholarship is allowed per 100 pupils in the school.

There are also two forms of Exhibitions. These are:—

a. Local Exhibitions to enable students to complete their education at some college or school where scientific instruction of an advanced character may be obtained. Grants of £25 per annum, for one, two, or three years are made for this purpose when the locality raises a like sum by voluntary subscriptions. And if the student attend a State School, such as the Royal School of Mines in London, the Royal College of Chemistry in London, or Royal College of Science in Ireland, the fees are remitted.

b. Royal Exhibitions of the value of £50 per annum tenable for three years, to the Royal School of Mines,

London, and the Royal College of Science, Dublin, are given in competition at the May examinations. Six are awarded each year—three to each institution. The exhibitions entitle the holder to free admissions to all the Lectures, and to the Chemical and Metallurgical Laboratories at those two institutions.

The competition for the Whitworth Scholarships, tenable for three years, is also in part determined by the results of the May examinations.

A grant in aid of a new building, or for the adaptation of an existing building, for a School of Science may be made at a rate not exceeding 2s. 6d. per square foot of internal area, up to a maximum of £500 for any one school, provided that certain conditions are complied with and that the school be built under the Public Libraries and Museums Act, or be built in connection with a School of Art, aided by a Department building grant.

A grant towards the purchase of fittings, apparatus, diagrams, &c., of 50 per cent of the cost of them is made to Science Schools. And where a school is furnished with a laboratory, properly fitted up, payments are made on account of students who during the year receive 25 lessons in laboratory practice.

Special extra grants in the form of capitation payments are made in fully organised Science Schools.

UNIVERSITY OF OXFORD.

Waynflete Professor of Chemistry.—W. Odling, M.A., F.R.S.

Professor of Mineralogy.—N. S. Maskelyne, M.A., F.R.S.

Every Student must reside in one or other of the Colleges or Halls, or in licensed lodgings, for a period of three years, passing at least two examinations in Arts, and one in either Mathematics, Natural Science, Law, Modern History, or Theology, when, if he obtain a first, second, or third class, he can take his B.A. Degree; if he do not gain such honour he has to pass a third examination in *Literis Humanioribus*.

Lectures by the Waynflete Professor on Mondays, and Thursdays at the Museum.

Lectures on Elementary Organic Chemistry will be given by the Aldrichian Demonstrator of Chemistry, Mr. W. W. Fisher, M.A., of Corpus Christi College, on Wednesdays and Saturdays, at 11 a.m.; and Lectures on Elementary Inorganic Chemistry by Mr. W. F. Donkin, M.A., of Magdalen College, on Tuesdays and Fridays, at 11 a.m.

The Laboratory of the University will be open from 10 a.m. to 4 p.m. daily, and instruction in Practical Chemistry will be given by the Aldrichian Demonstrator, and by Mr. John Watts, B.A., of Balliol College.

A Course of Practical Instruction in Organic Chemistry will be conducted by Mr. W. H. Pike, Ph.D.

The fee for students working in the Laboratory for three days in the week during the Term is £3; for students working every day, £5.

Scholarships of about the value of £75 are obtainable at Christ Church, Magdalen, and other colleges, by competitive examination in Natural Science.

More detailed information may be obtained from the University Calendar; from the professors; from E. Chapman, Esq., M.A., Frewin Hall; and from the Sub-Librarian in the Radcliffe Library or the Museum.

UNIVERSITY OF CAMBRIDGE.

Professor of Chemistry.—G. D. Liveing, M.A., F.R.S.

Jacksonian Professor of Natural and Experimental Philosophy.—J. Dewar, M.A., F.R.S.

Demonstrators.—J. W. Hicks, M.A., W. J. Sell, B.A., and H. J. H. Fenton, B.A.

The Student must enter at one of the Colleges, or as a Non-collegiate Student, and keep terms for three years by residence in the University. He must pass the previous

examination in Classics and Mathematics, which may be done in the first or second term of residence, or, through the Oxford and Cambridge Schools Examination Board, or through the Senior Local Examinations, before commencing residence. He may then proceed to take a Degree in Arts, either continuing mathematical and classical study, and passing the ordinary examinations for B.A., or going out in one of the Honour Triposes.

The scholarships, ranging in value from £20 to £80 a year, are chiefly given for mathematical and classical proficiency. Scholarships are given for Natural Science in Trinity, St. John's, St. Peter's, Clare, Christ's, Sidney, Pembroke, Caius, and Downing Colleges; the examinations being at Easter, and in June and October.

The Chemical Laboratory of the University is open daily for the use of the Students. The Demonstrators attend daily to give instructions.

Non-collegiate Students are allowed to attend certain of the College Lectures and all the Professors' Lectures, and have the same University status and privileges as the other Students. They are under the superintendence of the Rev. R. B. Somerset, Orford House, Cambridge, from whom further information may be obtained.

The following are the Lectures on Chemistry for the ensuing Academical Year:—

MICHAELMAS TERM, 1879.

General Course, by the Professor of Chemistry, on Mondays, Wednesdays, and Fridays, at 12 noon. Begin Oct. 13.

Spectroscopic Analysis, by the Professor of Chemistry, on Mondays, Wednesdays, and Fridays, at 1.30 p.m. Begin Oct. 17.

Analysis, by the Professor and the Demonstrators of Chemistry. Daily. Begin Oct. 13. Also at St. John's College, by Mr. Main. Begin Oct. 15. Also at Caius College, by Mr. Pattison Muir; begin Oct. 15. Also at Sidney College. Begin Oct. 14.

Metals, by Mr. Pattison Muir. Monday, Wednesday, and Friday, at 10. Begin Oct. 15.

Physical Chemistry, by the Jacksonian Professor, on Tuesdays, Thursdays, and Saturdays, at 12 noon. Begin Oct. 14.

Elementary Organic Chemistry, by Mr. Main, at St. John's College, on Tuesdays, Thursdays, and Saturdays, at 11 a.m. Begin Oct. 14.

Volumetric Analysis, by a Demonstrator, on Tuesdays, Thursdays, and Saturdays, at 10 a.m. Begin Oct. 14. Catechetical Lectures, by Mr. Lewis, at Downing College, on Mondays, Wednesdays, and Fridays, at 9 a.m. Begin Oct. 13.

Elementary Lectures on the more important Minerals, by the Deputy Professor of Mineralogy.

LENT TERM, 1880.

General Course continued, by the Professor of Chemistry, on Mondays, Wednesdays, and Fridays, at 12 noon. Begin Jan. 28.

Analysis, by the Professor or Demonstrators of Chemistry. Daily. Begin Jan. 19. Also at St. John's College. Begin Jan. 30. Also at Caius College Laboratory. Begin Jan. 31. Also at Sidney College Laboratory. Begin Jan. 30.

Organic Chemistry, by the Jacksonian Professor, on Tuesdays, Thursdays, and Saturdays, at 12 noon. Begin Jan. 29.

General Course, begun by Mr. Main, at St. John's Laboratory, on Tuesdays, Thursdays, and Saturdays, at 11 a.m. Begin Jan. 28.

Non-metallic Elements, by Mr. Pattison Muir, at Caius Laboratory, on Mondays, Wednesdays, and Fridays, at 10 a.m. Begin Feb. 4.

Catechetical Lectures, by Mr. Lewis, at Downing College, on Mondays, Wednesdays, and Fridays, at 9 a.m. Begin Jan. 30.

EASTER TERM, 1880.

Some Special Department, by the Professor of Chemistry, on Mondays and Fridays, at 12 noon. Begin April 14. Analysis, by the Professor or Demonstrators of Chemistry. Daily. Begin April 26. Also at St. John's College. Begin April 24. Also at Caius College. Begin April 25. Also at Sidney College Laboratory. Begin April 22.

Elementary Chemistry, by the Demonstrator of Chemistry, on Tuesdays, Thursdays, and Saturdays, at 12 noon. Begin April 13.

General Course concluded, by Mr. Main, at St. John's Laboratory, on Tuesdays, Thursdays, and Saturdays, at 12 noon. Begin April 24.

Elementary Organic Chemistry and Analysis, by Mr. Pattison Muir, at Caius Laboratory, on Mondays, Wednesdays, and Fridays, at 10 a.m. Begin April 26.

Catechetical Lectures, by Mr. Lewis, at Downing College, on Mondays, Wednesdays, and Fridays, at 9 a.m. Begin April 30.

Crystallography, by the Deputy Professor of Mineralogy.

UNIVERSITY OF DUBLIN.—TRINITY COLLEGE.

The Chemical Laboratory will re-open on October 1st, 1879.

The University Professor of Chemistry, Dr. Emerson Reynolds, F.R.S., assisted by the Demonstrator of Chemistry, Mr. Early, conducts the undermentioned Courses of Laboratory instruction:—

The First Course of Practical Chemistry.—Michaelmas Term:—Qualitative Analysis and the Use of the Spectroscope. Hilary Term: Volumetric and Simple Gravitimetric Analysis. Trinity Term: Organic Preparations and Analysis.

Students can also attend the Professor's Lectures on General Chemistry, and repeat most of the experiments performed in the Theatre.

This Course terminates on the last day of June.

The Second or Advanced Course includes instruction in the higher branches of Experimental and Analytical Chemistry, and in Methods of Research. Students who take out this Course are free to devote their chief attention to the study of special departments of Chemistry as applied to Arts and Industries.

Summer Course of Practical Chemistry for Medical Students.—This Course commences on the first Monday in April and terminates on the 30th of June following. Students experiment in the Laboratory from 2 to 4 o'clock on Tuesdays, Thursdays, and Saturdays.

Special Courses.—Students can enter at any time throughout the academic year for short terms of Laboratory instruction in Medical or Pharmaceutical Chemistry, or in the Methods of Analyses of Water, Air, &c., for Sanitary Purposes.

Lectures in Medical and Pharmaceutical Chemistry.—Dr. Reynolds lectures at 2 o'clock on Tuesdays, Thursdays, and Saturdays, from the 1st of November to the 31st of March following.

All the classes are open to extra-Academic Students.

KING'S COLLEGE.

(DEPARTMENT OF ENGINEERING AND APPLIED SCIENCE.)

Professor of Chemistry.—C. L. Bloxam, F.C.S.

Demonstrator of Practical Chemistry.—J. M. Thomson, F.C.S.

Assistant Demonstrator.—G. S. Johnson, F.C.S.

On Tuesday and Friday at 10.20 a.m. Students of the First Year are admitted to the Course of Theoretical and Applied Chemistry. The Course commences with a View of the Forces which concur to the production of Chemical Phenomena, after which the laws of Chemical Attraction are discussed, and the Non-metallic elements and their principal Compounds are described.

The Metals and their principal compounds are next examined, care being taken to point out the applications

of the Science to the Arts; and the processes of the different Manufactures, of Metallurgy, and of Domestic Economy, are explained and illustrated.

Examinations of the Class, both *visd voce* and by written papers, are held at interval during the course at the usual Lecture hour.

Second Year.—Students attend in the Laboratory twice a week, on Tuesday and Friday, at 10.20, and they go through a course of Manipulation in the most important operations of Chemistry, including the first steps of Analysis.

Any Student of this Department may be admitted to this Class at any period of his study on payment of an extra fee.

Experimental and Analytical Chemistry in the Laboratory.—The object of this Class is to afford to Students who are desirous of acquiring a knowledge of analysis, or of prosecuting original research, an opportunity of doing so under the superintendence of the Professor and Demonstrator; Students may enter, upon payment of extra Fees, at any time except during the vacation, and for a period of one, three, six, or nine months, as may best suit their convenience. The laboratory hours are from ten till four daily, except Saturday, on which day the hours are from ten till one.

In addition to the Laboratory Fee, each Student defrays the expenses of his own Experiments. The amount of this expense, which is comparatively trifling, is entirely under his own control.

Special hours and fees are arranged for the convenience of such Third Year Students as wish to study Analytical Chemistry.

Fees.—Chemistry per term, £3 3s. od.; per ann., £8 8s. od.; Practical Chemistry per term, £4 4s. od.; per ann., £8 8s. od.; Experimental and Analytical Chemistry—One Month (daily attendance), £4 4s. od.; Three Months (daily attendance), £10 10s. od.; Six Months (daily attendance), £18 18s. od.; Nine Months (daily attendance), £26 5s. od. A student taking a month's ticket may attend daily during 1 month, or 3 days a week during 2 months, or 2 days a week during 3 months.

Rules as to Admission of Students.

I. The Academical Year consists of Three terms: Michaelmas Term, from beginning of October to the week before Christmas; Lent Term, from the middle of January to the week before Easter; Easter Term, from Easter to the beginning of July.

II. The days fixed for the Admission of New Students in the Academical Year 1879-80, are Tuesday, September 30, Tuesday, January 13, and Wednesday, April 14.

UNIVERSITY COLLEGE.

FACULTY OF SCIENCE.

Chemistry.—Professor Williamson, Ph.D., F.R.S.

I. GENERAL COURSE.

Lectures daily (except Saturday) from 11 to 12 a.m., up to the last week in March.

Exercises on Tuesdays, Wednesdays, Thursdays, and Fridays, from 9 to 10 a.m.

Fees for the Course, £7 7s.; Perpetual, £9 9s.; for the Half Course, £4 4s.; for the Organic Course alone, £2 2s. Fee for the Exercise Class, £2 2s.

The instruction in this Class is of two kinds, consisting partly of Experimental Lectures by the Professor, partly of Exercises and personal instruction on the subject of the Lectures by an Assistant.

Attendance on the Exercise Class (conducted by Mr. Temple A. Orme, F.C.S.) enables Students to do their work more effectually and rapidly than they can do it by themselves.

A. *The first half* of the course, to Christmas, includes those parts of Chemistry which are required for the Matriculation Examination of the University of London.

B. The second half of the Course, from January to March, includes the following subjects:—

1. Preparation and properties of the chief *Metals*, including their characteristic reactions and most important salts. Detection of metallic poisons. Quantitative estimation of metals. Principles of classification. Monoatomic, diatomic metals, &c.

Organic Chemistry commences in the second week in February, and occupies five Lectures weekly till about the end of March. It includes a study of the characteristics and metamorphoses of the chief organic acids, bases, alcohols, ethers, colouring-matters, &c. Methods of ultimate and proximate analysis. Determination of molecular weights. Theory of types; of compound radicals. Phenomena of fermentation, &c.

Training of Teachers.

Teachers of Chemistry are trained in the theory and practice of their profession. A two years' Course is absolutely requisite for this purpose; but Students will with advantage devote a longer period to it.

The first year is occupied with attendance on the Courses of Chemistry and of Analytical Chemistry. In the second year the Student again attends the Course of Chemistry, and is intrusted with teaching-work in conjunction with the Tutors of the class. At the same time he continues to work in the Laboratory at analysis and original research.

In order to qualify themselves for rising to the higher ranks of the Profession, gentlemen remain for a further period, in which case they may obtain remunerative work in teaching through the recommendation of the Professor.

It must not, however, be supposed that a study of Chemistry alone, however complete, is sufficient to qualify a man to teach the Science effectively. A competent knowledge of Physics, Mathematics, and either French or German must necessarily be acquired at some period of the Student's Course.

II.—ANALYTICAL AND PRACTICAL CHEMISTRY.

A. Birkbeck Laboratory.

Assistants.—C. A. Bell, B.A., M.B., and Frank L. Teed.

When accompanied or preceded by attendance on the lectures on Chemistry, the Laboratory Course qualifies Students in the application of Chemistry to the Manufacturing Arts, Metallurgy, Medicine, or Agriculture, &c. Instruction is given in the principles and processes of gas-analysis.

The Laboratory and offices are open daily from 9 a.m. to 4 p.m., from the 3rd of October until the middle of July, with a short recess at Christmas and at Easter. Saturday, from 9 to 2.

Fees, for the Session, 25 guineas; six months, 18 guineas; three months, 10 guineas; one month, 4 guineas; exclusive of the expense of materials.

B.—SUMMER PRACTICAL COURSES.

Chemical Theatre.

1. *Elementary Course. (Men.)*—About Forty Lessons, of one hour each, commencing in the first week of May.

The first six weeks of the Course are occupied by the study of the chief non-metallic elements and their simple compounds. Metallic salts, &c., are subsequently studied.

Fees—including the cost of materials and apparatus: for the Course, £5 5s.; for a Second Course, £3 3s.

2. *Senior Course. (Men.)*—This Course consists of Twenty Lessons of two hours each, commencing in the first week in May.

The first half of the Course includes tests for fixed and volatile organic acids, nitrogenised acids, sugars, glycerin, alkaloids, &c.

The second half of the Course includes tests for mineral poisons in organic mixtures; also tests for organic bodies, such as alkaloids, when mixed with other organic substances.

Volumetric methods of the quantitative analysis of sugar and urea, chlorides, phosphates, hardness of water, alkalimetry, are practised.

Analysis of milk and ashes of blood.

Fees—including the costs of materials and apparatus: for the Course, £5 5s.; for a Second Course, £3 3s.

III.—SUMMER MATRICULATION COURSE. (Men.)

TEMPLE A. ORME, F.C.S.

This Course includes those parts of Chemistry which are required for the Matriculation Examination of the University of London.

The Course consists of about Twenty Lessons in Practical Chemistry, and of an equal number of oral lessons. These lessons will begin on April 13th, 1880, at 11.

Fee, including cost of materials and apparatus, £4 4s.

ELEMENTARY CHEMISTRY.

Chemical Theatre. (Women).

Lectures—Wednesday and Friday, from 4 to 5.

A Class of Elementary Chemistry, including the subjects required for Matriculation, will be given during the Winter Session by C. A. Bell, B.A., M.B., Chief Assistant in the Chemical Laboratory, commencing Wednesday, October 8.

The instruction will consist partly of Lectures, partly of Laboratory Experiments performed by the Students.

Fees for the Course, including use of apparatus and materials, £4 4s.

Chemical Technology.

Professor CHARLES GRAHAM, D.Sc.

Assistant.—Mr. Henry Brown.

The Course of instruction in this Department is designed to afford to Students who propose to devote themselves to industrial pursuits in which Chemistry plays an important part, or to prepare themselves for the profession of Consulting Chemist, the instruction essential for their success in their future line of work. It will also be found of great value in two of the branches (Organic and Inorganic Chemistry) in which the Degree of Doctor of Science can be taken at the University of London.

Assuming that the Student enters for a three year's study, the following will give an idea of the nature of the work during the period:—

In the first year the Student will attend Lectures on Theoretical Chemistry, and work at Analytical Chemistry in the Chemical Laboratory, and will also attend Lectures on Mathematics, Mechanics, and Physics. The Mechanical Drawing Class should also be attended during the first year by all Students.

In the second year, the Student will again attend the Lectures on Theoretical Chemistry, and will begin his study of Applied Chemistry by attendance on the Lectures in this subject, and by practical work in the Laboratory on the applications of Chemistry.

The third year will also be chiefly occupied with attendance on the Lectures on Chemical Technology, and in practical work connected therewith in the Laboratory.

In the second and third years the Student will, in addition to the foregoing subjects, which are common to all, attend Lectures and work at such other branches of Pure and Applied Science as may be deemed advisable after consultation with the Professor.

Students entering the College with more advanced scientific knowledge will be able to shorten the Course described to two years, or even one year, as may be found advisable.

For the convenience of those already engaged in business, and of those from other causes prevented from entering for a longer period of study, it is arranged that they can attend a Course of Lectures upon any one subject of Applied Chemistry without being required to attend any other lectures, either in Applied Chemistry, or in other subjects.

In the Session 1880-81, it is proposed to treat of the following subjects:—

- (1) The Chemistry of the Alkali trade.
- (2) Soap, Glass, Pottery, Cements.
- (3) Agricultural Chemistry.
- (4) Distilling, Vinegar-making, Bread- and Biscuit-making, or the Chemistry of Brewing.

Should a sufficient number of Students desire a Course of Lectures on some subject of Applied Chemistry other than those above mentioned, the Professor will give such either in lieu of, or in addition to, those mentioned.

Students desirous of working at subjects not included in the foregoing Courses, such as Photography and Photographic materials, Paper-making, Gas-tar products, the products of the Distillation of Wood, Tanning, and other Chemical industries, will receive individual instruction in the Laboratory.

Fees—for each Course, £2 2s.; for the four Courses together, £5 5s.

Chemical Laboratory.

The instruction in the Laboratory in Chemical Technology, will consist of the examination and valuation of raw materials used, and of the final products obtained, in various manufacturing industries, and of experimental examination of the processes employed in the arts and manufactures.

The Laboratories are open daily from 9 a.m. to 4 p.m., from the 2nd of October until the middle of July, with a short recess at Christmas and at Easter. Saturday, from 9 to 2.

Fees—for the Session, 25 guineas; six months, 18 guineas; three months, 10 guineas; one month, 4 guineas; exclusive of the expense of materials.

ROYAL SCHOOL OF MINES.

The mode of instruction at the Royal School of Mines is by systematic Courses of Lectures, by written and oral examinations, by practical teaching in the Laboratories, in the Drawing Office, and in the Field.

To become Associates at the Royal School of Mines it is necessary to pass through the following course of study:—

1st Year—Inorganic Chemistry, with practice in laboratory; Mechanical Drawing, both terms.

2nd Year—1st Term, Applied Mechanics; Physics, with practice in Laboratory. 2nd Term, Mineralogy; Mechanical Drawing, both terms.

3rd Year (Mining Division)—1st Term, Mining; Assaying. 2nd Term, Geology, with practice in laboratory and field. Metallurgical Division, 1st and 2nd Terms—Metallurgy, with practice in laboratory. Geological Division, 1st Term—Natural History, with practice in laboratory. 2nd Term—Geology, with practice in laboratory and field; Palaeontological Demonstrations.

The courses of instruction are distributed over three years, but those students who possess sufficient knowledge may, if they think fit, pass through the whole in two years, by presenting themselves during the current year for examination in the subjects allotted to the first and second years.

Students desirous of obtaining the distinction of Associate of the Royal School of Mines, who have already acquired a knowledge of the subjects of the first two years, may proceed at once to the courses of the third year, by passing the first class examinations in those subjects before the Professors of the Royal School of Mines, and paying a fee of £1 for each examination.

During the Session 1879-80 the following courses of Lectures will be delivered:—

40 Lectures on Inorganic Chemistry, commencing Oct. 6, 1879; 30 Lectures on Organic Chemistry, commencing Jan. 19, 1880, by E. Frankland, F.R.S.

80 Lectures on Biology, by T. H. Huxley, LL.D., F.R.S., commencing Oct. 6, 1879.

36 Lectures on Applied Mechanics, by T. M. Goodeve, M.A., commencing Oct. 6, 1879.

60 Lectures on Physics, by F. Guthrie, F.R.S., commencing Nov. 17, 1879.

80 Lectures on Metallurgy, by J. Percy, M.D., F.R.S., commencing Oct. 20, 1879.

60 Lectures on Mining, commencing Nov. 10, 1879, and 40 Lectures on Mineralogy, commencing Feb. 23, 1879, by W. W. Smyth, M.A., F.R.S.

50 Lectures on Geology, by John W. Judd, F.R.S., commencing Feb. 23, 1880.

20 Lectures on Mechanical Drawing, by J. H. Edgar, M.A., commencing Oct. 11, 1879.

The Lectures on Chemistry, Physics, Mechanics, Biology, and Geology are delivered at the Science Schools, South Kensington, where instruction is also given in the Chemical, Physical, Biological, and Geological Laboratories. The Lectures on Mineralogy, Mining, Metallurgy, and Mechanical Drawing are given at the School of Mines in Jermyn Street, where the Metallurgical Laboratory is situated.

The Laboratories for instruction in chemical manipulation, in qualitative and quantitative analysis, the technical application of analysis, and in the method of performing chemical researches, are under the direction of Dr. Frankland, and will be opened on Wednesday, October 1, 1879.

The charge for instruction in the Chemical Laboratory is £12 for three months, £9 for two months, and £5 for one month.

The Metallurgical Laboratory is conducted by Mr. R. Smith, under the direction of Dr. Percy, and is devoted to practical instruction in Metallurgy, especially in Assaying. The nature of this instruction will be adapted to the special requirements of the Student. It comprises:—Assaying in all its branches, especially of the more important metals, such as iron, copper, lead, tin, alloys of silver and gold, &c.; and the examination of ores and metallurgical products.

The ability of the Student to make trustworthy assays is in every case thoroughly tested; and no certificate of competency is given to a Student who has not furnished satisfactory proof that he is able to obtain accurate results.

The charge for instruction in the Metallurgical Laboratory is £15 for three months, £12 for two months, and £7 for one month.

Lectures to Working Men.—Short Courses of Lectures at suitable periods of the year are given in the evening to Working Men. These courses are systematic, and arranged so as to illustrate, within a period of two years, the principal subjects taught at the institution. Those for the ensuing Session include Geology, Metallurgy, Physics, and Biology.

The following Exhibitions, Scholarships, and Prizes are awarded in connection with the School:—

Nine Royal Exhibitions, each of £50 per annum, three of which are competed for annually at the May examinations of the Department of Science and Art.

Two Royal Scholarships of £15 each to the best students of the first year; and One Royal Scholarship of £25 to the best second year's student.

The Edward Forbes Medal and Prize for Biology and Palaeontology; the De la Beche Medal for Mining; and the Murchison Medal and Prize for Geology.

The public will be admitted to the lectures on payment of £4 for each course of 40 or more lectures, and £3 for the course of 30 and under 40 lectures.

The fee for students desirous of becoming Associates is £30 in one sum on entrance, or two annual payments of £20 each. Students presenting themselves for re-examination must pay a fee of £1 for each subject.

Persons who have taken either a first or a second class certificate in the advanced stage in any subject in science at the examinations held by the Science and Art Department, and who show that they are *bona fide* Science Teachers, may attend the Day Lectures gratuitously, provided that they be examined in at least one subject, paying fee for such examination of £1 per course.

Perpetual Tickets are issued, which entitle the holder to attend all present and future courses of lectures upon payment of £40.

Officers of the Army and Navy, Her Majesty's Consular and Diplomatic Officers, Officers of the Crown at home or on furlough, and acting Mine Agents and Managers of Mines, are admitted to the Lectures at half the above charges.

UNIVERSITY OF ABERDEEN.

Professor of Chemistry.—J. S. Brazier, F.C.S.

I. *Systematic Course.*—The Lectures are delivered on the first five days of each week during the College session. They commence with the discussion of the General Principles of Chemical Philosophy, including the Atomic Theory and the Chemical Relations of Heat. The Non-metallic and Metallic Elements and their Compounds are fully treated of, together with their more important applications to the Arts. The latter part of the course is devoted to the subject of Organic Chemistry. Examinations are held at fixed periods during the Session. The fee is £3 3s.

II. *Practical Course.*—This course is given during the Summer Session. It is chiefly devoted to practice in Qualitative Analysis, with the view of enabling the Student to test unknown substances, poisons, the animal secretions, &c. The fee is £3 3s.

III. *Laboratory Pupils.*—The Chemical Laboratory is open during the College Session on the first five days in each week, from 10 a.m. till 3 p.m. The course of instruction is under the direction of the Professor of Chemistry and of the Teaching Assistant.

UNIVERSITY COLLEGE, BRISTOL.

Professor of Chemistry.—E. A. Letts, Ph.D., F.R.S.E.

Lecturer.—W. W. J. Nicol, M.A.

Inorganic Chemistry.

Monday, Wednesday and Friday, 10 to 11.

This Course will be continued during the First and Second Terms, and will relate to Chemical Philosophy, the Chemistry of the Non-Metals, and the Chemistry of the Metals. Special attention will be given to the applications of Chemistry to the arts and manufactures. Tutorial Lectures will be given once a week by the Lecturer on those points in the course which require detailed explanation.

Fee for the Course, £4 4s.

For students who have paid Laboratory fees in the Session to the amount of £8 8s., this fee will be reduced to £3 3s.

Chemical Analysis.

A Course of about Twenty Lectures to be delivered during the First and Second Terms.

These lectures are recommended to all studying Practical Chemistry, and are intended to familiarise the student with the chief methods of Qualitative and Quantitative Analysis, and will partly take the form of demonstrations.

Organic Chemistry.

This Course will relate to the more important groups of the Carbon Compounds.

Fee, £2 2s. for this course alone; £5 5s. for this and the Winter Course together.

Practical Chemistry.—Laboratory Instruction.

The Laboratory will be open daily from 10 a.m. to 5 p.m., except on Saturdays, when it will close at 1 p.m. Instruction will be given in the Laboratory in all branches of Practical Chemistry, including Qualitative and Quantitative Inorganic and Organic Analysis, the preparation of Chemical Products, and Inorganic and Organic Research. Special facilities will be afforded to those who desire to study Practical Chemistry as applied to the different processes employed in the Arts and Manufactures.

Fees—

	6 Days a Week.	3 Days a Week.	2 Days a Week.
Per Session	18 gns.	10 gns.	7½ gns.
" Two Terms	13 "	7½ "	5½ "
" One Term	7 "	4 "	3 "
" Month	3 "	2 "	1½ "

In order that Students may have an opportunity of acquiring some knowledge of Applied Chemistry, excursions to some of the Mines and Manufactories of the neighbourhood will be made once a fortnight (on Saturdays) during the Summer Term. They will be conducted by the Professor or by the Lecturer. Past or present Students of the College desirous of taking part in these excursions are invited to apply to the Professor of Chemistry at the commencement of the Summer Term.

Evening Lectures.

Lecturer.—W. W. J. Nicol, M.A.

Tuesday and Thursday, 8 to 9.

This course will consist of Two Lectures a week; they will be devoted to the consideration of the Principles of Chemistry and the Study of the chief Non-Metallic Elements. In treating of the various products under the latter heading special attention will be devoted to their applications in the Arts and Manufactures.

Fee, 8s. per Two Terms.

Practical Class.

Professor.—E. A. Letts, Ph.D.

Lecturer.—W. W. J. Nicol, M.A.

Wednesday and Friday, 7 to 9.

A Practical Class will be formed for instruction in Qualitative and Quantitative Analysis provided a sufficient number of Students enrol. The Fee for the course (extending over the first and second terms) will be £4 4s.

Scholarships.

The following College Scholarships, open to women as well as men, will be competed for in October:—

One Chemical Scholarship of the value of £25, tenable for one year.

Three General Scholarships of the value of £25, £15, and £10 respectively, tenable for one year.

Four Scholarships for Women offered by the Clifton Association for Promoting the Higher Education of Women, will also be competed for. Of these, two will be entrance Scholarships; that is, they will be open only to those women who have not already commenced systematic study at the College. The other two will be open to all women excepting past and present scholars. Each of the four will be of the value of £15, tenable for one year, at University College, Bristol. Two of them may be augmented up to the maximum sum of £50 each, should such aid be necessary.

The Chemical Scholarship will be awarded principally by the marks obtained in Chemistry; but in case the best Candidates in it are nearly equal, account will be taken of their marks in other subjects.

ROYAL AGRICULTURAL COLLEGE, CIRENCESTER.

CHEMICAL DEPARTMENT.

Lectures and Laboratory Instruction are given in Organic, Inorganic, and Analytical Chemistry.

THE YORKSHIRE COLLEGE, LEEDS.

Professor of Chemistry.—T. E. Thorpe, Ph.D., F.R.S., F.C.S.

Lecture Courses.

1. General Course on Inorganic and Organic Chemistry—Monday, Tuesday, Wednesday, and Thursday, at 4 p.m., from October to the end of March. Fee for the Course, £4 4s.

2. Lectures on Laboratory Practice and Chemical Calculations—Thursday, at 10 a.m., during the First and Second Series. Fee, £1 1s.

3. Lectures on the Chemistry of the Non-Metals—Saturday, at 12 a.m., during the First and Second Terms. Fee, 10s. 6d.

Laboratory Courses.

The College Laboratory will be open daily from 9 a.m. to 1 p.m., and from 2 to 5 p.m., except on Saturdays, when it will close at 1 p.m.

Fees for the Session—Students working six days per week, £17 17s.; four, £13 13s.; three £11 11s.; two, £8 8s.; one, £4 4s.

Class in Practical Chemistry, Saturday mornings, from 9 to 12, during First and Second Series. Fee £1 11s. 6d.

Practical Chemistry for Medical Students.—On Monday and Wednesday, from 9 to 11 a.m., from May to the end of July.

Evening Classes.

A Course of twenty Lectures by Prof. T. E. Thorpe, F.R.S., on the Elements of Inorganic Chemistry (the Non-Metals) will begin during the first and second Terms, on Fridays, at 8 p.m., beginning October 17. Fee, 10s. 6d.

A Course of Twenty Lectures by Mr. C. H. Bothamley on the Elements of Inorganic Chemistry (the Non-Metals) will be given during the first and second Terms, on Mondays, at 8 p.m., beginning Oct. 13.

Scholarships.

The Cavendish Scholarship. Value £50 per annum, tenable for one year. Awarded for investigations made by the Candidates in any branch of Natural Science taught in the College.

The Salt Scholarship. Value £20 per annum, tenable for two years.

Akroyd Entrance Scholarships. Value £25 per annum, tenable for three years. Intended for the encouragement of the study of Natural Science. One of these Scholarships will be awarded annually, if in the opinion of the Examiners any Candidate shall possess sufficient merit.

The Clothworkers' Company Scholarships. Four Scholarships, each of the value of £25 per annum, and tenable for one year. Each Scholar will be required to attend regularly the Lectures and Courses of instruction given in the First and Second Years' Course of the Department of Textile Industries at the discretion of the Instructor.

UNIVERSITY OF DURHAM. COLLEGE OF PHYSICAL SCIENCE, NEWCASTLE.

Professor of Chemistry.—A. Freire-Marreco, M.A.
Demonstrator—J. T. Dunn, B.Sc.

Junior Division.—General Principles of Chemistry. History of the Non-Metallic Elements. History of the Metals and their more important Native and Artificial Compounds. Principles of Qualitative Analysis. Elements of Organic Chemistry. Senior Division.—Organic Chemistry. Elements of Applied Chemistry, including Chemical Mineralogy and Analysis.

Practical Chemistry.—The Laboratory is open from 10 a.m. to 1 p.m., and from 2 to 5 p.m., except on Saturdays, when it closes at 1 p.m. Laboratory Fees.—Students working six days per week, £5 5s. per term; alternate days, £3 3s.; one day per week, £1 1s.

Arrangements for Laboratory work in the evening and during vacation will be made.

The Session will commence on October 6.

Courses of Study.—Students will be distinguished into Regular and Occasional. Regular Students will be required to follow such a course of study in the subjects professed in the College as will enable them to pass the Examinations for the title of Associate in Physical Science.

Occasional Students will attend such classes as they may select. Every candidate for admission as a regular student must pass an examination on entrance, in reading, writing from dictation, English or Latin Grammar, arithmetic (including decimals), and geography. Registered students in medicine are exempted from this examination, or students who produce a certificate of having passed either of the two following examinations:—

1. Durham Senior Examination of Persons not Members of the University, held in June.

2. Durham Examination for Students in Arts in their first year, or any examination of a similar nature that may be accepted by the Council.

Associateship in Physical Science.—Every candidate for the Associateship in Physical Science, will be required to satisfy the examiners in three, at least, of the four subjects,—Mathematics, Physics, Chemistry, and Geology,—in an examination, to be held at the beginning of his second year.

The examination in Chemistry comprises:—General Principles of Chemistry. Elements of Inorganic Chemistry. Elements of Quantitative Analysis, including a Practical Examination.

The examination in Chemistry for Candidates at the end of their second year comprises:—Elements of Chemistry. Applied Chemistry. Advanced Qualitative Analysis, including a Practical Examination. Elements of Quantitative Analysis.

Exhibitions.—Three Exhibitions of £15 each will be awarded in October next, to Candidates desirous of attending the first year course of study in the College.

Candidates must pass the entrance examination, and will, in addition, be examined in the following subjects:—Algebra, up to Quadratic Equations. Euclid, Books I., II., and III. And one of the following special subjects, to be selected by the Candidate:—Geology.—Text Book: Page's "Introductory Text Book." Heat.—Text Book: "Orme on Heat." Chemistry.—Text Book: Gill's "Chemistry for Schools," omitting chapters xiii., xvi., xx., and xxi. Natural History.—Text Book: Nicholson's "Elementary Text Book of Zoology," or Oliver's Elementary Botany, Part I.

Candidates must send their names to the Secretary, on or before the 27th of September, and specify, at the same time, the special subject in which they desire to be examined.

The examination will be held at the College, and will commence on Monday, the 6th October.

Two similar Exhibitions will be awarded at the next examination of "Persons not members of the University," which will be held at Durham, and elsewhere, in June next, to those candidates who shall most distinguish themselves in subjects allied to Physical Science.

Scholarships.

T. Y. Hall Scholarship.—This Scholarship, of the yearly value of £20, tenable for three years by students attending two or more of the classes, will be awarded on the result of the first examination for the Associateship in Science.

Charles Mather Scholarship.—This Scholarship, of the yearly value of about £40, will be awarded on the result of the Final Examination for the Associateship in Science, and is tenable for one year from the time of obtaining the Associateship in Science, provided the Scholar continues his studies in the College to the satisfaction of the Professors.

Nathaniel Clark Scholarship.—This Scholarship, of the value of £15 for one year, will be awarded in October to that student who shall pass the First Examination for the Associateship in Science, and who shall be most distinguished in Chemistry and Geology. The Scholar will be required to attend the classes of Chemistry and Geology, so as to be qualified to take those subjects for the Final Examination for the Associateship in June next.

OWENS COLLEGE, MANCHESTER.

Professor and Director of the Chemical and Metallurgical Laboratories.—H. E. Roscoe, B.A., Ph.D., F.R.S., F.C.S.

Professor of Organic Chemistry.—C. Schorlemmer, F.R.S.

Demonstrators and Assistant Lecturers.—Mr. W. C. Williams, F.C.S., Mr. Thomas Carnelley, D.Sc., and Mr. J. B. Hannay, F.C.S.

Lecture Courses.

Systematic Chemistry.—*Junior Class.*—Tuesday, Thursday, and Saturday, from 9.30 to 10.30 a.m., during Michaelmas and Lent Terms. Comprising—(1) The laws of Chemical Combination; (2) a description of the physical and chemical properties and the mode of preparation of the Non-Metallic Elements and of their Compounds.

Senior Class.—Monday, Wednesday, and Friday, from 9.30 to 10.30 a.m., during the Michaelmas and Lent Terms, comprising—(1) The Chemistry of the Metals and of their most important Compounds; (2) Organic Chemistry.

The instruction in Systematic Chemistry is given by means (a) of Experimental Lectures and (b) of Tutorial Classes.

Fee.—For each Class, £2 12s. 6d.; for both Classes, £4 14s. 6d.

A Tutorial Class, meeting in Sections, will also be held, which all members of the Junior and Senior Classes will be required to attend, unless specially exempted by the Principal and the Professor. Extra fee for this Class, 10s. 6d. This fee is not included in the composition fees payable by regular Students.

Organic Chemistry.—Professor C. Schorlemmer, F.R.S., Monday, Wednesday, and Friday, from 10.30 to 11.30 a.m.

General Course (from October to the end of March).—The subject of this course is the Chemistry of the Carbon Compounds, wherein the branch of Organic Chemistry is more fully and completely treated than in the general course in Systematic Chemistry.

Extended Course (from the beginning of April to the end of the Session).—This course is suited to the requirements of students preparing for the B.Sc. examination and for those who have previously attended the general course and wish to become more intimately acquainted with the subject. The course will treat of the History of the Old and New Theories, and of the most recent important discoveries in Organic Chemistry, &c.

Fee for the General Course, £2 12s. 6d.; for the Extended Course, £1 11s. 6d.; for both Courses, £3 10s.

Chemical Philosophy.—Prof. C. Schorlemmer, F.R.S., Saturday, from 9.30 to 10.30 a.m.

Sketch of the History of Chemistry; Development of Modern Chemistry; Chemical Law and Theories; Relation of Chemistry to Physics.

Fee, £1 11s. 6d.

Technological Chemistry.—Dr. Carnelley will give in the Michaelmas and Lent Terms a Course of about Twenty Lectures, on Mondays, from 2.30 to 3.30 p.m., on Water, Air, and the Chemistry of Fuel and the Gas Manufacture.

Fee, £1 1s.

Analytical Chemistry.—Mr. W. C. Williams, F.C.S., Thursday, from 10.30 to 11.30 a.m.

This Course will treat of the methods of Qualitative and Quantitative Analysis, and is intended to supplement the instruction in Practical Chemistry.

Fee, £1 11s. 6d.

Analytical and Practical Chemistry.

LABORATORY COURSES.

The Chemical Laboratories will be open for Students daily from 9.30 a.m. until 4.30 p.m., except on Saturdays, when they will be closed at 12.30 p.m.

Fees for the Session.—For six days per week, £21; for four days per week, £17 17s.; for three days per week, £13 13s. Students entering the Laboratory Class at or after Christmas will be charged two-thirds of the fees for the whole Session.

Fees for shorter periods.—For six months, £17 17s.; for five months, £15 15s.; for four months, £13 13s.; for three months, £10 10s.; for two months, £7 7s.; for one month, £4 4s. Students entering under this scale are entitled to work on every day during the week.

The Metallurgical Laboratory will be open daily during the same hours as those of the Chemical Laboratories, for instruction in Practical Assaying and the examination of Ores and Metallurgical products.

Fees the same as those for the Chemical Laboratory Course.

A Course of Lectures on Metallurgy will probably be given in the course of the session if a sufficient number of students offer themselves.

Entrance Exhibitions.

I. Victoria Exhibition (Classics), £15.

II. Wellington Exhibition (Greek Testament), £15.

III. Dalton Mathematical Exhibition, £15.

The Victoria and Dalton Exhibitions are renewable for a second year.

IV. Grammar School Scholarships, £15 per annum, tenable for three years; open to scholars of the Manchester Grammar School only.

V. Two Oxford and Two Cambridge Local Exhibitions, giving free admission to lecture classes in the College for one year, and renewable for two years further are awarded annually on the results of the Oxford and Cambridge Examinations held in Manchester.

VI. Gilchrist Scholarship, £50 per annum, tenable for three years; awarded on the results of the Matriculation Examination of the University of London, in June, 1879.

VII. Rumney Scholarships, £45 per annum, tenable for three years. The next competition will take place in 1880.

VIII. Ramsbottom Scholarship, £40 per annum, tenable for two years. The next competition will take place in 1880.

IX. Crace-Calvert Scholarships, £25 per annum, tenable for two years. The next competition will take place in June, 1880. This scholarship is open only to duly qualified members of the Evening Chemistry Classes.

Fellowship.

The Langton Fellowship, £150 per annum, tenable for three years. Candidates must have been students in the College for not less than three sessions, and must during their studentship or within one year after the close of the same have obtained a degree of some University of the United Kingdom, or been elected to the Associateship of the College. The next competition for the Fellowship will take place in 1881.

Scholarships.

The following (except the Shakspeare Scholarship) are open to the competition of students of the College only.

I. Victoria Scholarship (Classics), £40 per annum, tenable for two years.

II. Wellington Scholarship (Greek Testament), £40 per annum, tenable for two years.

III. Shuttleworth (Political Economy), £50, tenable for one year.

IV. Shakspeare Scholarship (English Language and Literature), £40 per annum, tenable for two years.

V. Bradford History Scholarship, £45 per annum, tenable for one year, and renewable for a second year.

VI. Dalton Chemical Scholarships, two, each of £50 per annum, tenable for two years.

VII. Dalton Mathematical Scholarships, one Senior and one Junior Scholarship, of the value of £25 each, tenable for one year.

VIII. Platt Scholarships (Physiology), two, each of £50 per annum, tenable for two years.

IX. Heginbottom Physical Scholarship, £20 per annum, tenable for two years.

X. Ashbury Scholarships (Engineering), two, each of £25 per annum, tenable for two years.

Prizes.

I. Lee Greek Testament Prizes, one of £25 and one of £12 10s. value.

II. Classical Prizes, value £5. Junior Classical Prizes, value £5 and £2 10s.

III. Shuttleworth History Prize, value £5.

IV. English Essay and Poem Prizes, each of the value of £5.

V. Early English Text Society's Prizes.—A selection of the Society's publications offered to the competition of students in the Day and in the Evening Classes respectively.

VI. New Shakspeare Society's Book Prizes.

VII. Bryce Law Prize, value £10.

VIII. Cobden Club Book Prizes (Political Economy).

IX. Dalton Natural History Prize, value £15.

X. Engineering Essay Prize, value £5.

ROYAL COLLEGE OF SCIENCE FOR IRELAND,
STEPHEN'S GREEN, DUBLIN.

Professor of Practical and Theoretical Chemistry.—W. Noel Hartley, F.C.S.

The Chemical and Metallurgical Laboratories, under the direction of Mr. Hartley, are open every week-day during the Session, except Saturday. Instruction is given in the different branches of Analytical Chemistry, including Assaying, and in the methods for performing Chemical Research. Fee, for the Session of nine months, £12; or for three months, £5; or for one month, £2.

There are four Royal Scholarships of the value of £50 each yearly, with Free Education, including Laboratory Instruction, tenable for two years; two become vacant each year; they are given to Students who have been a year in the College. There are also nine Exhibitions attached to the College, of the yearly value of £50 each, with Free Education, including Laboratory Instruction, tenable for three years; three become vacant each year.

A Diploma of Associate of the College is granted at the end of the three years' course.

ANDERSON'S COLLEGE, GLASGOW.

Professor of Chemistry.—William Clittmar, F.R.S.E.

Chief Assistant.—M. T. Buchanan.

Lecture Assistant.—Robert Lennox.

Junior Assistants.—James M. Bowie and George A. Barling.

A Course of 100 Experimental Lectures on Chemistry: Daily, Saturdays excepted, from 10 to 11, commencing about the end of October. The Lectures up to the end of the year are devoted to the elements of Chemical Philosophy and to the Chemistry of the Non-metallic Elements. After the new year the Course divides into two branches, viz., the Chemistry of the Metals (on the Mondays and Tuesdays) and Organic Chemistry, select chapters (on the Wednesdays, Thursdays, and Fridays). Six written examinations are held during the Session, which all the members of the class are required to attend.

Fee, £2 2s.

A Course of Tutorial Lessons in connection with the above Lectures will be given by one of the Assistants. Free to the members of the class.

The Laboratory for Practical Instruction in all branches of analysis, including technical assaying, and for original research is open daily (Saturdays excepted) during the Winter Session from 10 to 5, during Summer from 9 to 5. The teaching is conducted on the tutorial system, each student working by himself and on his own subject. The

Laboratory is furnished with all the necessities for chemical investigation.

Fee for the Winter Session, £10 10s.; Summer Session, £6 6s.; two sessions, if paid in advance, £15 15s., or £2 2s. per month.

THE

"YOUNG" CHAIR OF TECHNICAL CHEMISTRY,
ANDERSON'S COLLEGE.

Professor.—Edmund J. Mills, D.Sc. (Lond.), F.R.S.

Senior Assistant.—Mr. J. Snodgrass.

Junior Assistant.—Mr. J. Bicket.

This Chair has for its object the instruction of Students in Chemistry as applied to the various branches of industry in Chemical and other works, Metallurgy, Agriculture, &c.

LECTURES.—*Principal Course.*—A Course of Fifty Lectures will be delivered on Mondays, Tuesdays, and Wednesdays, at 10 a.m., commencing on November 3rd. The Lectures will be illustrated with Experiments, Diagrams, and Models, as well as by the actual inspection of Manufacturing Processes; and the progress of the Students will be tested by periodical Examinations. The earlier Lectures will have reference to units of weight and measure, to the calculations necessitated by Chemical operations, and to the nature and laws both of the Chemical process and its results. A particular subject will then be considered in comparatively minute detail, embracing for this session Coal-Gas: the concluding lectures will have special reference to Lubricating Oils.

Fee for the Course, Two Guineas. To Laboratory Students, One Guinea.

Subsidiary Course.—A Course of Thirty Lectures will be delivered on Mondays, Tuesdays, and Wednesdays, at 9 a.m., commencing in May. These Lectures are more particularly intended for Dyers, Colour Manufacturers, Brewers and Distillers, Tar Rectifiers, Drysalterns, and others interested in a knowledge of Technical Organic Chemistry.

Fee for the Course, Two Guineas.

Technical Physical Chemistry.—Mr. J. Snodgrass, Senior Assistant, has arranged to deliver a series of Thirty Lectures and Demonstrations on Chemical Apparatus.

Fee for the Course, Half-a-Guinea. Laboratory Students, Half-a-crown.

Laboratories.—The Laboratories are open daily from 10 to 4, and on Saturday from 10 to 1 o'clock for practical working by the Students, under the superintendence of the Professor and his Assistants.

The Fee for attending the Laboratories is £20 per Session of Nine Months, £14 10s. for Six Months, £7 10s. for Three Months.

Students must have a fair acquaintance with elementary Chemistry.

The New Laboratory Buildings, immediately contiguous to the site formerly occupied, are now erected and in order for occupation. They comprise four stories, with a lecture room in the rear, and are exclusively devoted to the purposes of this Chair.

The Trustees, having had under consideration the requirements of Inventors, Patentees, and others whose investigations require isolation and privacy, have included in the arrangements Five Private Laboratories, which will, it is expected, meet a demand hitherto unsupplied in this country.

Library.—A Students' Library Society was founded in 1875. Its objects are to provide a collection of standard chemical works, and to maintain a regular supply of chemical journals. A large number of works have already been purchased or bestowed, and nine journals are received. Annual subscription, Half-a-crown.

Memorandum as to Bursaries.

The Trustees of the "Young" Chair have the superintendence of the Bursaries—regulating the appointment and terms on which they shall continue to be held.

The Nominees of Donors to be appointed if they pass the necessary examinations.

The Bursaries are of the amount of £50 each per annum, tenable for three years, during which the Bursars shall be required to give their whole time and attention to the Lectures and Laboratory duties of the "Young" Chair, paying the ordinary fees. Candidates to have attained sixteen years of age on application, to be of good moral character, and to pass such examinations as may be prescribed by the Trustees in the ordinary branches of an English education and the elementary principles of Chemistry. The Bursaries to be liable to forfeiture on the Bursars failing to exhibit approved progress under the Professor of the Chair, or being guilty of conduct, in the opinion of the Trustees, unworthy of their position. The Bursaries are only given to those whose means are limited, and who intend following some branch of Manufacturing Chemistry.

CHEMICAL LECTURES, CLASSES, AND LABORATORY INSTRUCTION.

CRYSTAL PALACE COMPANY'S SCHOOL OF ART, SCIENCE, AND LITERATURE. SCHOOL OF PRACTICAL ENGINEERING. *Principal*—Mr. J. W. Wilson, Assoc. Inst. C.E.—This school was established with the purpose of affording to Students of Civil or of Mechanical Engineering the advantage of thorough practical instruction in the rudiments of either profession, and in the manipulation of materials. The leading object is to prepare Students, by systematic practical instruction, for professional articles, so that on entering an Engineer's office or works the pupil may at once be useful to his Principal, and enabled to take advantage of the opportunities for learning open to him, because he has mastered the elementary details of the profession. The school is also available for Students already articulated, who desire instruction either in the offices or shops. The Colonial Section is designed particularly for gentlemen who are going to the Colonies or abroad, as explorers or settlers. The object proposed is to afford them so much practical knowledge of scientific and mechanical work and expedients as shall enable them best to utilise the means at their disposal, especially when entirely dependent on their own resources.

Special Courses of Lectures are delivered by independent Lecturers not on the Staff of the School, and are an addendum to the curriculum. These Special Courses include a Course of Six Lectures on the Chemistry of Manufactures and Mines; and a Course of Six Lectures on the Mechanics of Practical Mining.

LADIES' DIVISION.—The School was established to utilise the valuable Courts and Collections of the Crystal Palace for the purposes of instruction in Art, Science, &c., so that education of the highest class might be afforded on reasonable terms under most advantageous conditions. The system of tuition is, for some subjects, in the manner of private tutorial instruction by the best masters, but other subjects are taught on the University method, in accordance with the regulations laid down by the Syndicate of the University of Cambridge, by whom some of the lectures and classes are conducted. A student may take lessons in one or several studies at option. The School is a centre for both the University of Oxford and the University of Cambridge Local Examinations, the Oxford Examination for Women, and for the Cambridge Higher Local Examination. The following examinations will be held in the Ladies' Division during 1879-80:—Cambridge Local, December, 1879; Oxford Local and Oxford Examination, for Women, May, 1880; Cambridge Higher Examination, June 1880. The session opens on October 1.

The Courses of Lectures during the ensuing term will include one on National Geography, on Mondays, by Prof. H. G. Seeley, F.R.S. Fees: Twelve Lectures and Classes, £1 1s. Twelve Lectures, 10s. 6d. One Lecture, 1s.

BERNERS COLLEGE OF CHEMISTRY AND THE EXPERIMENTAL SCIENCES, 44, Berners Street, W.—Prof. E. V. Gardner, F.A.S., M.S.A. The Laboratory is open morning and evening throughout the year.

BIRKBECK LITERARY AND SCIENTIFIC INSTITUTION, Southampton Buildings, Chancery Lane.—Inorganic Chemistry. Lectures:—Elementary, Tuesdays, 8.30 to 9.30; Advanced, Saturdays, 7 to 8. Practice:—Elementary, Saturdays, 4 to 6; Advanced, Saturdays, 8 to 10. Teacher, Geo. Chaloner, F.C.S. Organic Chemistry:—Course of Thirty Lectures will be given on Tuesday evenings, at 7 o'clock, by Mr. H. Chapman Jones, F.C.S., commencing on October 7th. Practical Organic Chemistry:—This Class will meet in the Laboratory of the Institution, under Mr. Chapman Jones's direction, on Saturdays from 4 to 6 and from 8 to 10 p.m.

NEW CENTRAL SCHOOL OF CHEMISTRY AND PHARMACY, 173, Marylebone Road, London.—Mr. A. P. Luff, F.C.S., and Mr. J. Woodland, M.P.S.

ONSLow COLLEGE OF SCIENCE, Pond Place, Fulham Road, S.W.—Special Evening Classes in Inorganic and Organic Chemistry, &c. The Chemical Laboratory is open on Friday Evenings from 6 to 10 p.m. and on Saturday from 2 to 10 p.m.

ROYAL VETERINARY COLLEGE, Camden Town.—Professor of Chemistry, Mr. R. V. Tuson.

SCHOOL OF PHARMACY OF THE PHARMACEUTICAL SOCIETY OF GREAT BRITAIN, 17, Bloomsbury Square.—The school opens on Wednesday, the 1st of October. Lectures on Chemistry and Pharmacy, by Professor Redwood, on Monday, Tuesday, and Wednesday mornings, at 9 a.m. The Laboratories for Practical Instruction in Chemistry as applied to Pharmacy, &c., under the direction of Prof. Attfield, will be open daily at 10 a.m. throughout the Session. They are fitted up with every convenience for the study of the principles of Chemistry by personal experiment, synthetical as well as analytical. They are specially designed for the student of Pharmacy, but are also well adapted for the acquirements of a knowledge of Chemistry in its application to Manufactures, Analysis, and Original Research. There is no general class for simultaneous instruction, each student following an independent course of study always determined by his previous knowledge; pupils can therefore enter for any period at any date. We have in previous Students' Numbers drawn attention to the fact that the Professors strongly advise all learners, both before and during attendance at the school, to avoid studying merely by way of "preparation for examination." The one desire should be education, that is knowledge accompanied by enlightenment of the understanding, mental training, mental discipline, and general elevation of the intellect. As for "examination," students of the School may rest assured that if they, guided by the Professors, work diligently, thoughtfully, deliberately, and thoroughly, they will with ease and pleasure pass any examination in the subjects in which they have thus been truly educated.

Council Prizes.—At the end of each of the five months' Courses of Lectures on Chemistry and Pharmacy, and Botany and Materia Medica, a Bronze Medal and Certificates of Merit, and at the close of the Session (ten months) a Silver Medal and Certificates of Honour and Merit, are offered for competition by the Council. In the Class of Practical Chemistry, the Silver Medal, two Bronze Medals, and Certificates of Honour and Merit, offered by the Council, are competed for at the end of the Session only.

SOUTH LONDON SCHOOL OF CHEMISTRY, 325, Kennington Road.—Dr. John Muter, F.C.S. Daily, at 10 a.m. Sixty Lectures on Theoretical Chemistry, and Junior and Senior Course of Practical Chemistry.

THE WESTMINSTER COLLEGE OF CHEMISTRY AND PHARMACY, Lambeth Road, S.E.—Messrs. Wills and Wootton. Daily, at 9.30 a.m. Theoretical and Practical Chemistry.

WORKING MEN'S COLLEGE, 91, Blackfriars' Road.—Teacher of Chemistry, Mr. J. D. Allen.

CHEMICAL LECTURES AT LONDON HOSPITALS.

Chemical Schools and Colleges.	WINTER SESSION.				SUMMER SESSION.			
	Lecturers on Chemistry.	Days and Hours.	Fees.		Lecturers on Chemistry.	Days and Hours.	Fees.	
			One Course.	Perpetual.			One Course.	Perpetual.
St. Bartholomew's Hosp. and College	Dr. Russell, F.R.S.	M. W. F., 9	6 16/6	9 9	Dr. Russell	M. Tu. F., 11	3 3	£ s.
Charing Cross Hospital and College	Mr. Heaton	M. W. F., 11	5 5		Mr. Heaton	M. W. Th. F.	3 3	
St. George's Hospital ..	Mr. Wanklyn	Tu. Th. S., 11 1/2	7 7		Mr. Wanklyn	M. W. F., 10	4 4	
Guy's Hospital	Dr. Debus, F.R.S., and Dr. Stevenson	Tu. Th. S., 11 1/2	7 7		Dr. Debus	M. W. F., 10 to 1	7 7	
King's College and Hosp.	Dr. Stevenson	M. W. Th., 10 1/2	8 8	11 11	Mr. Bloxam and Mr. Hartley, & Mr. Thomson	M. W. Th., 10 1/2	6 6	8 8
London Hospital	Mr. Bloxam, F.C.S., Mr. Thomson, and Mr. Johnson	M. Tu. W. Th., 11	7 7	7 7	Mr. Thomson	M. Th. S., 9	2 2	
St. Mary's Hospital ..	Dr. Tidy	M. W. Th., 4	6 16/6		Dr. Tidy	W. F. S.	4 4	
Middlesex Hospital ..	Dr. Wright	M. W. Th. Fr., 9	6 6	8 8	Dr. Wright	M. W. F., 3	3 3	
St. Thomas's Hosp. & Schl.	Mr. W. Foster	Tu. Th. F., 10 7	7 7		Mr. W. Foster	M. Th. F., 10	6 6	
University Col. & Hosp.	Dr. Bernays	Daily (ex. S.), 11 7	7 7	9 9	Dr. Bernays	Daily (ex. S.), 11	4 4	7 7
Westminster Hospital ..	Dr. Williamson, F.R.S. and Dr. Graham	W. Th. F., 3	6 6		Dr. Williamson & Dr. Graham	M. W. F., 10	4 4	
	Dr. Dupré, F.R.S.				Dr. Dupré			

BIRMINGHAM.—MIDLAND INSTITUTE.—Mr. C. J. Woodward, B.Sc. Tuesday and Thursday, at 8 p.m.; Friday, at 7; and Saturday, at 3.

BIRMINGHAM.—QUEEN'S COLLEGE.—Mr. A. Bostock Hill, M.D.

BRISTOL MEDICAL SCHOOL.—Mr. T. Coomber, F.C.S.

LIVERPOOL ROYAL INFIRMARY SCHOOL OF MEDICINE.—J. Campbell Brown, D.Sc. Lond., F.C.S.

SCHOOL OF TECHNICAL CHEMISTRY, 7 and 9, Hackin's Hey, Liverpool.—Mr. A. Norman Tate.

COLLEGE OF CHEMISTRY, LIVERPOOL.—Mr. S. H. Johnson, F.C.S.

LEEDS SCHOOL OF MEDICINE.—Prof. T. E. Thorpe, Ph.D., F.R.S.

LEEDS MECHANICS' INSTITUTION.—Mr. G. Ward, F.C.S.

MANCHESTER GRAMMAR SCHOOL.—Mr. Francis Jones, F.C.S., F.R.S.E.

MANCHESTER MECHANICS' INSTITUTION.—Mr. M. A. Watts, M.A.

QUEENWOOD COLLEGE, near Stockbridge, Hants.—Mr. E. W. Prevost, Ph.D., F.C.S., F.R.S.E.

SALFORD WORKING MEN'S COLLEGE, EVENING CLASSES.—Teacher of Chemistry, Mr. G. H. Hurst.

SHEFFIELD BOROUGH ANALYSTS' LABORATORY, 1 and 3, Surrey Street.—Mr. A. H. Allen, F.C.S. Day and Evening Classes.

SHEFFIELD SCHOOL OF MEDICINE.—Mr. A. H. Allen, F.C.S. A course of forty-five Lectures on Inorganic and Organic Chemistry is delivered by Mr. Alfred H. Allen during the Winter session. The Summer Course of Practical Chemistry is under the direction of Mr. A. H. Allen.

ABERDEEN SCHOOL OF SCIENCE AND ART MECHANICS' INSTITUTION.—Mr. Thomas Jamieson, F.C.S.

DUNDEE LITERARY INSTITUTION CHEMICAL AND PHYSICAL LABORATORY.—Lecturer on Chemistry, Mr. Frank W. Young, F.C.S., and Mr. John Thomson.

UNIVERSITY OF EDINBURGH.—Prof. A. Crum Brown, F.R.S.E.

SCHOOL OF MEDICINE, EDINBURGH.—Dr. Stevenson Macadam, F.R.S.E., Mr. Falconer King, and Mr. Ivison Macadam.

SCHOOL OF PHARMACY AND CHEMISTRY, EDINBURGH.—The instruction qualifies for graduation in Medicine and Science in the University of Edinburgh and other Examining Boards. Chemistry, Mr. Drinkwater, F.C.S., L.R.C.P.; Materia Medica and Pharmacy, Dr. Urquhart; Botany, Mr. D. M'Alpine, B.Sc. Lond. Day and Evening Classes.

MINTO HOUSE MEDICAL SCHOOL, CHAMBERS STREET, EDINBURGH.—Lectures and Classes.

GLASGOW UNIVERSITY.—Prof. J. Ferguson.

GLASGOW MECHANICS' INSTITUTION.—Mr. R. R. Tatlock, F.R.S.E., F.C.S.

GLASGOW VETERINARY COLLEGE.—Mr. Stephen Cooke, F.C.S.

SCHOOL OF CHEMISTRY, 138, Bath Street, Glasgow.—Dr. Wallace, Mr. Tatlock, and Dr. Clark. Day and Evening Classes.

CHEMICAL LABORATORY, 144, West Regent Street, Glasgow.—Dr. Milne. Day and Evening Classes.

ANALYTICAL LABORATORY, 88, Hope Street, Glasgow.—Dr. A. T. Machattie, F.C.S. Day and Evening Classes.

QUEEN'S COLLEGE, BELFAST.—Dr. Andrews, F.R.S., &c.

QUEEN'S COLLEGE, CORK.—Dr. Maxwell Simpson.

QUEEN'S COLLEGE, GALWAY.—Dr. T. H. Rowney.

ROYAL COLLEGE OF SURGEONS IN IRELAND.—Dr. C. A. Cameron.

DUBLIN, CARMICHAEL SCHOOL.—Dr. C. R. C. Tichborne.

DUBLIN, CATHOLIC UNIVERSITY.—Mr. Campbell.

DUBLIN, DR. STEVENS'S HOSPITAL AND MEDICAL COLLEGE.—Mr. McHugh.

ON TWO NEW ELEMENTS IN ERBIA.

By P. T. CLÈVE.

TOWARDS the end of last year M. Marignac discovered in erbia, which, up to that time, had been considered as the oxide of a single metal, erbium, the oxide of a new element very strongly characterised, viz., ytterbia. A short time afterwards M. Nilson found in erbia another oxide, scandia, the salts of which are as colourless as those of ytterbia. The substance which gives to the salts of erbia their red colouration and magnificent absorption spectrum, that is to say true erbia, is still unknown. I resolved to extract if possible from the old erbia this colouring principle. I had at my disposal a considerable quantity of material, almost entirely free from ytterbia. M. Nilson

was so good as to give me the residues of the extraction of scandia and ytterbia. I found, however, even after hundreds of separations, that it was impossible to obtain a red oxide of constant molecular weight. I was therefore led to suppose the presence of another unknown oxide, and I asked M. Thalén to examine the absorption spectrum of the fraction that I regarded as pure erbia, and at the same time to compare its spectrum with the spectra of the residues rich in ytterbia and yttria. Some absorption bands in the last fractions suggested the idea that the colour of erbia is occasioned by the presence of three oxides in the absorption spectrum. I then re-united the reddest fractions, the molecular weight of which was from 126 to 127 (RO), and submitted them to a long series of decompositions, treating one fraction (A) for ytterbia, another (B) for yttria, and an intermediary third, in which true erbia ought to be concentrated. At the same time I tried to concentrate the colouring-matter in the residues rich in ytterbia (A) and in yttria (B). When I had exhausted the material at my disposal I asked M. Thalén to kindly examine the five fractions. This examination led to the following results:—

For the examination of absorption spectra of the five given fractions, the liquid was enclosed in a glass trough with parallel sides of a thickness of 0.008 m., or in test-tubes of about 0.010 m. diameter. In order to obtain sufficient, but very different dispersions, I employed for each of the fractions, sometimes two, sometimes six Flint prisms, whose angle of refraction was 60°. The registration is referred to the solar spectrum, and the positions of the absorption bands relate to Angström's normal spectrum of the sun. The figures below express the wave-lengths in 100,000ths.

The absorption bands common to all the fractions, which bands must probably be attributed to erbia, are as follow:—

Colour of Rays.	Wave-Length.	Remarks.
Red	6660—6680	Feeble
	6515—6545	Strong
	6475—6515	Tolerably strong
Yellow	5400—5415	" "
	5225—5235	Very strong
Green	5185—5225	Strong
	4865—4877	Strong
Indigo	4475—4515	Tolerably strong

A very marked difference is, however, observed between the following bands, according as they are examined with one or other of the liquids in question.

	Length of Wave.	Fraction A.		Erbium (?) Mean Fractions (126—127).	Fraction B.	
		From Residues of Ytterbia.	Erbia (126—127).		Erbia.	From Residues rich in Yttrium.
x	6840	Strong	Tolerably strong	None	None	None
y	6400—6425	None or trace only	Trace	Feeble	Feeble	Very strong
z	5360	None	None or trace	Trace	Feeble	Tolerably strong

It is then seen that the band *x* belongs to the fractions situated near to ytterbia, whilst it does not exist in the fractions derived from yttria. But it is quite the contrary with respect to the bands *y* and *z*; indeed, these bands, which are wanting altogether in the residues of ytterbia, show themselves more and more clearly as we approach yttria.

It follows from these researches that the spectrum of old erbia must be attributed to three distinct oxides. In fact, the colour of the solution of the different fractions materially differs. Whilst the fractions treated for ytterbia are of a rose colour with a tinge of violet, the fractions treated for yttria have an orange tinge. Although I possess considerable quantities of the mixture of these three oxides, I am convinced that it would be useless to continue these researches until I have been able to procure still larger quantities. For the radical of the oxide

placed between ytterbia and erbia, which is characterised by the band *x* in the red part of the spectrum, I propose the name of *Thulium*, derived from Thulé, the ancient name of Scandinavia. The atomic weight of this metal must be about 113 (its oxide being RO); at least its oxide is found concentrated in the fractions which have for their molecular weight 129. The atomic weight of true erbia, to which the common bands are to be attributed, is probably from 110 to 111. Its oxide is of a light rose colour.

The third metal characterised by the bands *y* and *z*, and which is found between erbia and terbia, must have a lower atomic weight than 108. Its oxide appears to be yellow; at least, all the fractions of a molecular weight lower than 126 are more or less yellow. I propose for this metal the name of *Holmium*, Ho, derived from the latinized name of Stockholm, in the neighbourhood of which so many minerals rich in yttria are to be found.

It only remains for me to express my gratitude to M. Thalén for the trouble he has taken in all these researches.

Referring to the above communication of M. Clève, Prof. J. Lawrence Smith made the following observations:—

I have latterly had the opportunity of conversing with several of the *savants* who are occupying themselves in the study of the earths of the yttrium and cerium group. Notwithstanding the interesting results they have already obtained, they still have more or less doubt as to the clearness of those results, and the conclusion which may be deduced from them on account of the difficulty of separating these earths one from the other.

Attention ought to be directed as much as possible towards the purification of the earths, as until they can be obtained pure only their relative position among the elements can be indicated. How far the impurities modify the absorption rays, whether in the luminous or in the ultra-violet part of the spectrum, is still an open question. The earth with which I have specially occupied myself—oxide of mosandrum—does not give absorption rays, and I am obliged to study it chemically, and I think I have obtained it in a tolerably pure state, but not sufficiently so to satisfy me. I may say that it is more insoluble in sulphate of potash than the terbia of Mosander. I have already indicated, moreover, some other special properties. With respect to the absorption rays of earths, I should call the attention of the Academy to a note by M. Lecoq de Boisbaudran and myself (*Comptes Rendus*, vol. lxxxviii., p. 1167, in which we have shown how in

an easy and complete manner some of the rays of didymium can be changed.—*Comptes Rendus*, September 1, 1879.

NOTES ON THE ALKALOIDS. A NEW TEST FOR PAPAVERINE.

By J. TATTERSALL.

PAPPAVERINE when brought in contact with concentrated sulphuric acid gives a light pink-violet colour. On heating this either disappears completely, or becomes a light grey, but the following reaction is more permanent and more characteristic of the alkaloids:—

Place the substance to be tested for papaverine in an evaporating basin, add a few drops of concentrated H₂SO₄,

and warm till complete solution is effected. Now add a fragment of sodium arsenate, and warm again over a small flame, inclining the dish to obtain as large a surface of liquid as possible. The original colour once more makes its appearance, but on continued application of heat it becomes cherry-red, and finally, as vapours of sulphuric acid begin to escape, dark bluish-violet. This colouration is very stable. When the contents of the dish are quite cold, add about 10 c.c. of water, and pour the orange liquid obtained into a flask, dilute once more, and add caustic soda to strongly alkaline reaction; the liquid rapidly darkens in colour, and when an excess has been added appears almost black; it is violet-red by reflection, and a pink straw colour by transmitted light. The alkaloids strychnine, brucine, morphine, salicine, atropine, narcotine, narceine, digitaline, picrotoxine, curarine, colchicine, and cantharidine do not exhibit this reaction, becoming, on subsequent addition of alkali, light orange or dirty yellow.

Codeine, when heated with concentrated H_2SO_4 and Na_3AsO_4 , gives a fine deep blue colour, much darker than the one produced by Fe_2Cl_6 under similar circumstances. On addition of water and alkali this becomes orange, and is characteristic of the alkaloid.

Hayfield Printing Co., Hayfield.

ON RADIANT MATTER.*

By WILLIAM CROOKES, F.R.S.

(Concluded from page 107.)

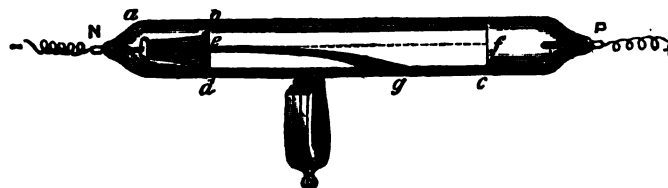
Radiant Matter is deflected by a Magnet.

I now pass to another property of Radiant Matter. This long glass tube (Fig. 14), is very highly exhausted; it has

This action of the magnet is very curious, and if carefully followed up will elucidate other properties of Radiant Matter. Here (Fig. 15) is an exactly similar tube, but having at one end a small potash tube, which if heated will slightly injure the vacuum. I turn on the induction current, and you see the ray of Radiant Matter tracing its trajectory in a curved line along the screen, under the influence of the horse-shoe magnet beneath. Observe the shape of the curve. The molecules shot from the negative pole may be likened to a discharge of iron bullets from a mitrailleuse, and the magnet beneath will represent the earth curving the trajectory of the shot by gravitation. Here on this luminous screen you see the curved trajectory of the shot accurately traced. Now suppose the deflecting force to remain constant, the curve traced by the projectile varies with the velocity. If I put more powder in the gun the velocity will be greater and the trajectory flatter, and if I interpose a denser resisting medium between the gun and the target, I diminish the velocity of the shot, and thereby cause it to move in a greater curve and come to the ground sooner. I cannot well increase before you the velocity of my stream of radiant molecules by putting more powder in my battery, but I will try and make them suffer greater resistance in their flight from one end of the tube to the other. I heat the caustic potash with a spirit-lamp and so throw in a trace more gas. Instantly the stream of Radiant Matter responds. Its velocity is impeded, the magnetism has longer time on which to act on the individual molecules, the trajectory gets more and more curved, until, instead of shooting nearly to the end of the tube, my molecular bullets fall to the bottom before they have got more than half-way.

It is of great interest to ascertain whether the law governing the magnetic deflection of the trajectory of

FIG. 14.



a negative pole at one end (a) and a long phosphorescent screen (b, c) down the centre of the tube. In front of the negative pole is a plate of mica (b, d) with a hole (e) in it, and the result is, when I turn on the current, a line of phosphorescent light (e, f) is projected along the whole length of the tube. I now place beneath the tube a powerful horseshoe magnet: observe how the line of light

Radiant Matter is the same as has been found to hold good at a lower vacuum. The experiments I have just shown you were with a very high vacuum. Here is a tube with a low vacuum (Fig. 16). When I turn on the induction spark, it passes as a narrow line of violet light joining the two poles. Underneath I have a powerful electro-magnet. I make contact with the magnet, and the

FIG. 15.



(e, g) becomes curved under the magnetic influence waving about like a flexible wand as I move the magnet to and fro.

line of light dips in the centre towards the magnet. I reverse the poles, and the line is driven up to the top of the tube. Notice the difference between the two phenomena. Here the action is temporary. The dip takes place under

* A Lecture delivered to the British Association for the Advancement of Science, at Sheffield, Friday, August 22, 1879

the magnetic influence; the line of discharge then rises and pursues its path to the positive pole. In the high exhaustion, however, after the stream of Radiant Matter had dipped to the magnet it did not recover itself, but continued its path in the altered direction.

By means of this little wheel, skilfully constructed by Mr. Gimingham, I am able to show the magnetic deflection in the electric lantern. The apparatus is shown in this diagram (Fig. 17). The negative pole (*a, b*) is in the form of a very shallow cup. In front of the cup is a mica screen (*c, d*), wide enough to intercept the Radiant Matter coming from the negative pole. Behind this screen is a mica wheel (*e, f*) with a series of vanes, making a sort of paddle-wheel. So arranged, the molecular rays from the

I have mentioned that the molecules of the Radiant Matter discharged from the negative pole are negatively electrified. It is probable that their velocity is owing to the mutual repulsion between the similarly electrified pole and the molecules. In less high vacua, such as you saw a few minutes ago (Fig. 16), the discharge passes from one pole to another, carrying an electric current, as if it were a flexible wire. Now it is of great interest to ascertain if the stream of Radiant Matter from the negative pole also carries a current. Here (Fig. 18) is an apparatus which will decide the question at once. The tube contains two negative terminals (*a, b*) close together at one end, and one positive terminal (*c*) at the other. This enables me to send two streams of Radiant Matter side by

FIG. 16.

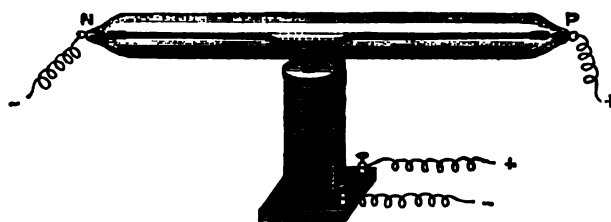


FIG. 17.

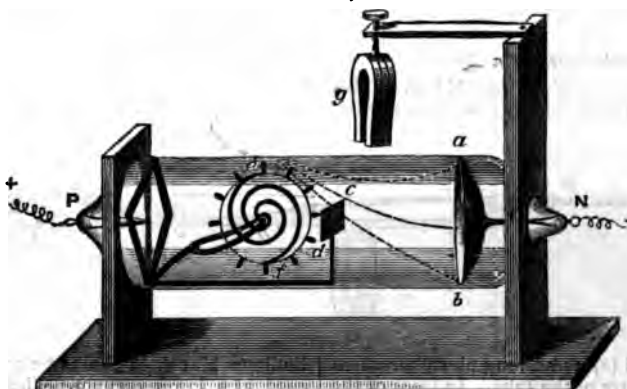
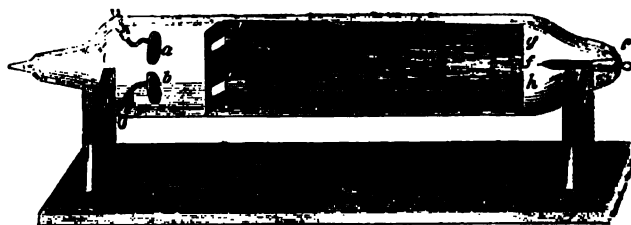


FIG. 18.



pole *a b* will be cut off from the wheel, and will not produce any movement. I now put a magnet, *g*, over the tube, so as to deflect the stream over or under the obstacle *c d*, and the result will be rapid motion in one or the other direction, according to the way the magnet is turned. I throw the image of the apparatus on the screen. The spiral lines painted on the wheel show which way it turns. I arrange the magnet to draw the molecular stream so as to beat against the upper vanes, and the wheel revolves rapidly as if it were an over-shot water-wheel. I turn the magnet so as to drive the Radiant Matter underneath; the wheel slackens speed, stops, and then begins to rotate the other way, like an under-shot water-wheel. This can be repeated as often as I reverse the position of the magnet.

side along the phosphorescent screen,—or by disconnecting one negative pole, only one stream.

If the streams of Radiant Matter carry an electric current they will act like two parallel conducting wires and attract one another; but if they are simply built up of negatively electrified molecules they will repel each other.

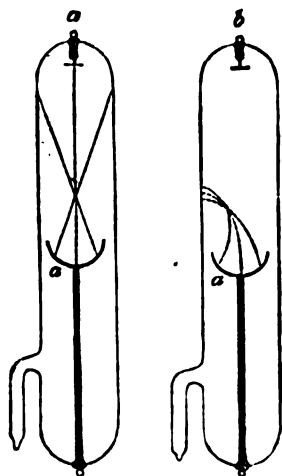
I will first connect the upper negative pole (*a*) with the coil, and you see the ray shooting along the line *d, f*. I now bring the lower negative pole (*b*) into play, and another line (*e, h*) darts along the screen. But notice the way the first line behaves; it jumps up from its first position, *d, f*, to *d, g*, showing that it is repelled, and if time permitted I could show you that the lower ray is also deflected from

its normal direction: therefore the two parallel streams of Radiant Matter exert mutual repulsion, acting not like current carriers, but merely as similarly electrified bodies.

Radiant Matter produces heat when its motion is arrested.

During these experiments another property of Radiant Matter has made itself evident, although I have not yet drawn attention to it. The glass gets very warm where the green phosphorescence is strongest. The molecular focus on the tube, which we saw earlier in the evening

FIG. 19.



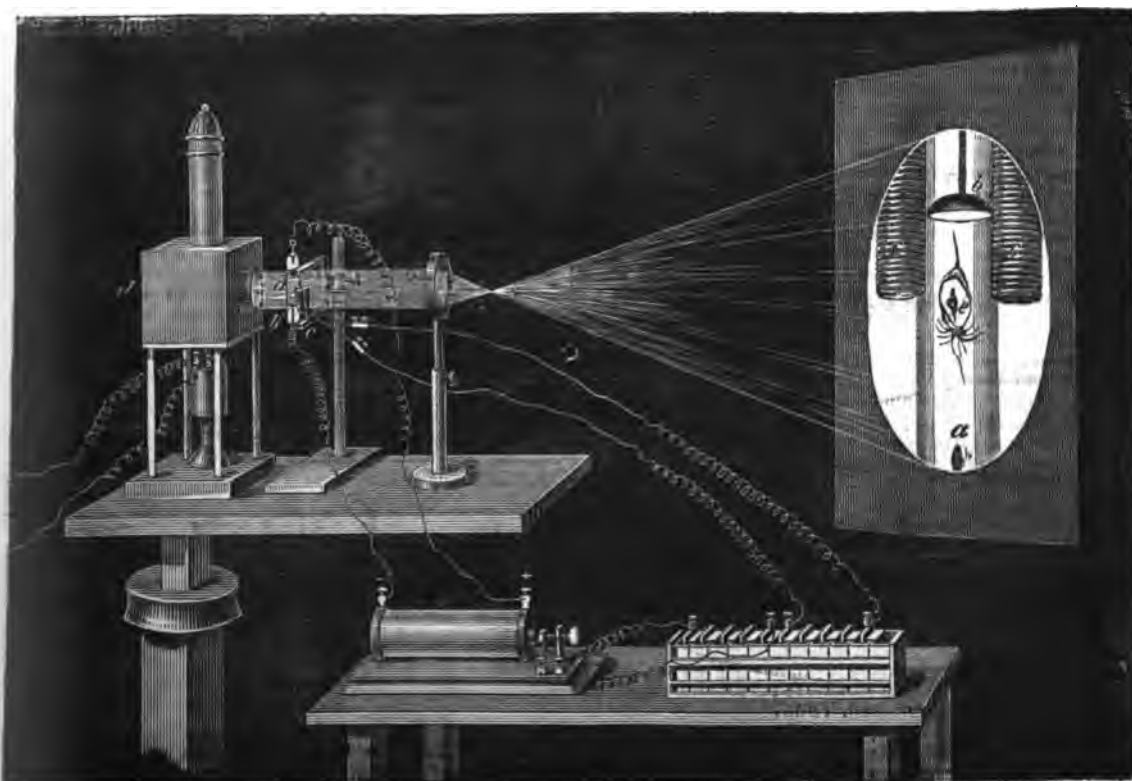
(Fig. 8) is intensely hot, and I have prepared an apparatus by which this heat at the focus can be rendered apparent to all present.

I have here a small tube (Fig. 19, a) with a cup-shaped negative pole. This cup projects the rays to a focus in the middle of the tube. At the side of the tube is a small electro-magnet, which I can set in action by touching a key, and the focus is then drawn to the side of the glass tube (Fig. 19, b). To show the first action of the heat I have coated the tube with wax. I will put the apparatus in front of the electric lantern (Fig. 20, d), and throw a magnified image of the tube on the screen. The coil is now at work, and the focus of molecular rays is projected along the tube. I turn the magnetism on, and draw the focus to the side of the glass. The first thing you see is a small circular patch melted in the coating of wax. The glass soon begins to disintegrate, and cracks are shooting starwise from the centre of heat. The glass is softening. Now the atmospheric pressure forces it in, and now it melts. A hole (c) is perforated in the middle, the air rushes in, and the experiment is at an end.

I can render this focal heat more evident if I allow it to play on a piece of metal. This bulb (Fig. 21) is furnished with a negative pole in the form of a cup (a). The rays will therefore be projected to a focus on a piece of iridio-platinum (b) supported in the centre of the bulb.

I first turn on the induction-coil slightly, so as not to bring out its full power. The focus is now playing on the metal, raising it to a white-heat. I bring a small magnet near, and you see I can deflect the focus of heat just as I did the luminous focus in the other tube. By shifting the magnet I can drive the focus up and down, or draw it completely away from the metal, and leave it non-luminous. I withdraw the magnet, and let the molecules have full play again; the metal is now white-hot. I

FIG. 20.



increase the intensity of the spark. The iridio-platinum glows with almost insupportable brilliancy, and at last melts.

FIG. 21.



The Chemistry of Radiant Matter.

As might be expected, the chemical distinctions between one kind of Radiant Matter and another at these high exhaustions are difficult to recognise. The physical properties I have been elucidating seem to be common to all matter at this low density. Whether the gas originally under experiment be hydrogen, carbonic acid, or atmospheric air, the phenomena of phosphorescence, shadows, magnetic deflection, &c., are identical, only they commence at different pressures. Other facts, however, show that at this low density the molecules retain their chemical characteristics. Thus by introducing into the tubes appropriate absorbents of residual gas, I can see that chemical attraction goes on long after the attenuation has reached the best stage for showing the phenomena now under illustration, and I am able by this means to carry the exhaustion to much higher degrees than I can get by mere pumping. Working with aqueous vapour I can use phosphoric anhydride as an absorbent; with carbonic acid, potash; with hydrogen, palladium; and with oxygen, carbon, and then potash. The highest vacuum I have yet succeeded in obtaining has been the 1-20,000,000th of an atmosphere, a degree which may be better understood if I say that it corresponds to about the hundredth of an inch in a barometric column three miles high.

It may be objected that it is hardly consistent to attach primary importance to the presence of *Matter*, when I have taken extraordinary pains to remove as much Matter as possible from these bulbs and these tubes, and have succeeded so far as to leave only about the one-millionth of an atmosphere in them. At its ordinary pressure the atmosphere is not very dense, and its recognition as a constituent of the world of *Matter* is quite a modern notion. It would seem that when divided by a million, so little *Matter* will necessarily be left that we may justifiably neglect the trifling residue and apply the term *vacuum* to space from which the air has been so nearly removed. To do so, however, would be a great error, attributable to our limited faculties being unable to grasp high numbers. It is generally taken for granted that when a number is divided by a million the quotient

must necessarily be small, whereas it may happen that the original number is so large that its division by a million seems to make little impression on it. According to the best authorities, a bulb of the size of the one before you (13.5 centimetres in diameter) contains more than 1,000,000,000,000,000,000,000,000 (a quadrillion) molecules. Now, when exhausted to a millionth of an atmosphere we shall still have a trillion molecules left in the bulb—a number quite sufficient to justify me in speaking of the residue as *Matter*.

To suggest some idea of this vast number I take the exhausted bulb, and perforate it by a spark from the induction coil. The spark produces a hole of microscopical fineness, yet sufficient to allow molecules to penetrate and to destroy the vacuum. The inrush of air impinges against the vanes and sets them rotating after the manner of a windmill. Let us suppose the molecules to be of such a size that at every second of time a hundred millions could enter. How long, think you, would it take for this small vessel to get full of air? An hour? A day? A year? A century? Nay, almost an eternity! A time so enormous that imagination itself cannot grasp the reality. Supposing this exhausted glass bulb, indurated with indestructibility, had been pierced at the birth of the solar system; supposing it to have been present when the earth was without form and void; supposing it to have borne witness to all the stupendous changes evolved during the full cycles of geologic time, to have seen the first living creature appear, and the last man disappear; supposing it to survive until the fulfilment of the mathematician's prediction that the Sun, the source of energy, four million centuries from its formation will ultimately become a burnt-out cinder; supposing all this,—at the rate of filling I have just described, 100 million molecules a second—this little bulb even then would scarcely have admitted its full quadrillion of molecules.†

But what will you say if I tell you that all these molecules, this quadrillion of molecules, will enter through the microscopic hole before you leave this room? The hole being unaltered in size, the number of molecules undiminished, this apparent paradox can only be explained by again supposing the size of the molecules to be diminished almost infinitely—so that instead of entering at the rate of 100 millions every second, they troop in at a rate of something like 300 trillions a second. I have done the sum, but figures when they mount so high cease to have any meaning, and such calculations are as futile as trying to count the drops in the ocean.

In studying this Fourth state of *Matter* we seem at length to have within our grasp and obedient to our control the little indivisible particles which with good warrant are supposed to constitute the physical basis of the universe. We have seen that in some of its properties Radiant Matter is as material as this table, whilst in other properties it almost assumes the character of Radiant Energy. We have actually touched the border land where Matter and Force seem to merge into one another, the shadowy realm between Known and Unknown which for me has always had peculiar temptations. I venture to think that the greatest scientific problems of the future

* The possible duration of the Sun from formation to extinction has been variously estimated by different authorities, at from 18 million years to 400 million years. For the purpose of this illustration I have taken the highest estimate.

† According to Mr. Johnstone Stoney (*Philosophical Magazine*, vol. 36, p. 141), 1 c.c. of air contains about 1000,000,000,000,000 molecules. Therefore a bulb 13.5 centims. diameter contains $13.5^3 \times 0.5236 \times 1000,000,000,000,000$ or 1,288,232,350,000,000,000 molecules of air at the ordinary pressure. Therefore the bulb when exhausted to the millionth of an atmosphere contains 1,288,232,350,000,000,000 molecules, having 1,288,231,061,747,650,000,000,000 molecules to enter through the perforation. At the rate of 100,000,000 molecules a second, the time required for them all to enter will be

1288,231,061,747,650 seconds, or
214,706,110,291,775 minutes, or
3,578,435,170,295 hours, or
149,099,380,092 days, or
411,258,321 years.

will find their solution in this Border Land, and even beyond; here, it seems to me, lie Ultimate Realities, subtle, far-reaching, wonderful.

"Yet all these were, when no Man did them know,
Yet have from wisest Ages hidden been;
And later Times things more unknown shall show.
Why then should witless Man so much misweene,
That nothing is, but that which he hath seen?"

NEW PROCESS FOR THE RAPID ESTIMATION OF PURE SUGAR IN RAW AND REFINED COMMERCIAL SUGARS.*

By P. CASAMAJOR.

(Concluded from page 108.)

Correction for Variations of Temperature.—When we operate at temperatures different from 15° C. or 60° F., the corrections can be made by using either of the tables of Gay-Lussac or those for the instrument of Tralles, which we owe to Prof. MacCulloch. I will, however, state that results that are sufficiently accurate may be obtained by proceeding as follows:—

For solutions which give, by the alcohometer from 77 to 70, we may multiply the excess of temperature above 60° F. by 0.2, and deduct the product from the reading of the alcohometer. For degrees of the alcohometer between 70 and 60, we may multiply this excess by 0.205, and for indications below 60° of the alcohometer, the excess of temperature above 60° F., must be multiplied by 0.21, and the product deducted. If a centigrade thermometer is used, the excess of temperature over 15° should be multiplied by 0.36 for solutions above 70° of the alcohometer. For solutions between 60° and 70° (alcohometer), the excess over 15° C. should be multiplied by 0.37, and for those below 60°, this excess should be multiplied by 0.38. In all cases, the product should be deducted from the direct reading of the alcohometer. The results obtained in this way do not differ materially from those given by the tables.

There is another correction for variation of temperature, which relates to the volume of standard solution to be taken for a weight of sugar equal to 19.8 gr. As the temperature rises, a greater volume of standard solution should be taken. If we use a Fahrenheit thermometer we take—

At 60° F.		50.0	c.c. of standard solution.
" 70		50.3	" " "
" 80		50.6	" " "
" 90		51.0	" " "
" 100		51.3	" " "

If we use centigrade thermometer, we should take—

At 15° C.		50.0	c.c. of standard solution.
" 20		50.25	" " "
" 25		50.5	" " "
" 30		50.8	" " "
" 35		51.2	" " "
" 40		51.4	" " "

If it should be preferred to use always the same volume of 50 c.c. of standard solution, we may take the following weights for the temperatures of a centigrade thermometer:

At 15° C.		19.8	grammes.
" 20		19.7	"
" 25		19.6	"
" 30		19.5	"
" 35		19.4	"
" 40		19.3	"

In the following table may be found the same data concerning methylic alcohol of various strengths, saturated with sugar, as are given in table No. 1 for ethylic alcohol.

* Read before the American Chemical Society, June 5, 1879.

TABLE No. 4.

Degrees of the alcohometer before saturation.	Ditto after saturation with sugar.	Degree of the saccharometer (Ventzke).	Grms. of sugar in 100 c.c.
92.5	91.8	17	0.44
83.5	77.1	13.2	3.43
82.7	76.5	—	—
81.5	75.0	—	—

From this table we may learn that the solution, saturated with sugar, standing at 77.1 by the alcohometer, should, when placed in the saccharometer tube, stand at 13.2. We also learn that the same saturated solution must contain 3.43 grammes of sugar in 100 c.c. This is a useful guide for the quantity of extra powdered sugar that should be added to 1 litre of methylic alcohol of 83.5°.

Method of Procedure in Testing.—The sugar to be tested should not be weighed before everything is ready, as otherwise it may dry out, after being weighed, and give too high a result. The mortar and pestle should be clean and dry, the cylinder should be filled with the standard solution to a line indicating 50 c.c., and a clean dry funnel should be ready with a paper filter in it. The weight of sugar to be taken is 19.8 grammes. This is transferred to the mortar as soon as weighed, and the standard solution should be poured in the mortar, and the sugars then ground with a heavy pestle until all lumps and large crystals are broken up.†

The mortar I have used for grinding sugar in presence of the standard solution has a capacity of 180 c.c., and the pestle is of such a size that it displaces 80 c.c. in this mortar.

After all the lumps have been broken up the contents of the mortar are poured on a filter standing over the cylinder, which previously held the standard solution. As there was some of this solution left in the cylinder, about 10 c.c. of the filtered solution is allowed to run into the cylinder, after which the funnel is taken from the cylinder and held over the mortar long enough to shake up the liquid in the cylinder and pour it back on the filter. After this the funnel is placed again over the cylinder, the contents of the mortar are poured again on the filter, and from 30 to 35 c.c. are allowed to run through the filter, which are generally sufficient for a test. In this filtered solution are placed, in succession, an alcohometer and a thermometer. To the alcohometric degree, corrected for temperature, is added the difference between 100 and the alcohometric degree of the standard solution. This sum represents the percentage of sugar.

Example.—The standard solution gives with the alcohometer 76.8 when corrected for temperature. The same alcohometer placed in the filtered solution marks 71°, while a centigrade thermometer shows a temperature of 28°. According to what has been already said, we take the difference between 28 and 15 = 13, and multiply this

* From this table we may also learn that if we apply methylic alcohol to Payen's process, we need have no fear of leaving in the sugar a small quantity of the solution made by saturating with sugar methylic alcohol at 92.5°. With ethylic alcohol, cane sugars are always left with too much impurity. Dr. Behr informs me that on this account he has never been able to obtain good results by this process. It is certainly worth while trying methylic alcohol. Cane sugars could be washed out with our standard solution, testing 77.1 (alcohometer), and the last particle of this could be removed by methylic alcohol of 92.5, saturated with sugar. The small quantity of this latter remaining after filtration would leave too little sugar to cause an appreciable error. The funnel with auxiliary vertical tube, of which I gave a description several years ago, is very well adapted to washing sugars out with alcohol, as it leaves them quite dry without the help of a filter pump (see *American Chemist*, Oct., 1875, vol. vi., p. 122, and *CHEMICAL NEWS*, vol. xxxii., No. 829, p. 183).

† Persons who test raw centrifugal sugars by this process must very soon see what *artificially coloured sugars* consist in. This has been a vexatious question, occupying the attention of eminent chemists. By washing sugar crystals with alcohol we may easily see that in the grains of some of them have been incorporated brown insoluble particles. In some cases these look like the scum from the defecation of the juice, and, in other samples, the colouring matter seems to be clay or some brown pigment. This colouring matter seems generally to have been added in a very clumsy manner, as after washing the crystals in alcohol it may be seen in lumps, speckles, the crystals with dark dots.

difference by 0.36, the product being 4.68. This quantity subtracted from 71 gives 66.32. The difference between 100 and 76.8 being 23.2, this quantity is added to 66.32, and the sum, 89.52, is the quantity of pure sugar in the sugar tested.

It has been suggested that the evaporation of methylic alcohol during the manipulations that have been described must interfere with the accuracy of the results obtained, but this does not seem to be the case. It might be possible to avoid evaporation entirely by leaving out the operation of grinding the mixture of sugar and methylic alcohol in a mortar; but, in this case, the results obtained would be less accurate on account of the impurities left in the sugar. Beyond avoiding working in a draft, no precaution has been taken to prevent evaporation. One reason why the evaporation of the alcohol seems to have so little effect on the result is that the operation of grinding and filtering occupy only a few minutes, during which no appreciable loss takes place.

NOTICES OF BOOKS.

Catechism of Agricultural Chemistry and Geology. By the late J. F. W. JOHNSTON. An entirely new edition, revised and enlarged by C. A. CAMERON. 78th thousand. Blackwoods, 1879.

To revise an old established favourite like this little Catechism is no easy task. It is hard for an editor to leave out statements which may not precisely commend themselves to his judgment, but which he does not know how to replace by anything more certain. It is hard to keep to the old limits and yet to add the more important facts discovered by recent researches. It is hard to maintain in the new part a style which came quite naturally to the original author. To say that Dr. Cameron has succeeded perfectly in an operation where failure is so common would be incorrect. But the new editor has undoubtedly improved the book, as it has been issued since Dr. Voelcker's revision of the original text in 1863. He ought to have done more, however. The old tables on pages 34, 36, and 37 needed revision; those on page 53 should have been replaced by sets of figures representing the results of properly conducted field-experiments: it is of no service to any one to tell them that a certain number (8) of extra bushels of wheat-grain were obtained from an acre of land by the use of 4 cwt. of "wheat manure," unless the experiment were tried in duplicate at least, the composition of the manure given, and all the other conditions of the experiment, natural and artificial, duly recorded and discussed. And even then such information and such details cannot be profitably used by the young learners, for whom the little work under notice is intended. Figures drawn from a much wider range, as well as higher class of experimental trials, are necessary for them. We hope Dr. Cameron will go through the book again before the issue of the 79th thousand, and make a good many more corrections and additions. With the exception of one of the old tables having been expanded into six, and a few figures and statements having been slightly changed, we do not see any great revision or enlargement of the present edition.

A Systematic Course of Practical Qualitative Analysis. By T. ELTOFT. Pp. vi. and 62. London: Simpkin, Marshall, and Co., 1879.

ALTHOUGH this pamphlet does not differ widely from other digests of qualitative chemical operations, it bears throughout distinct marks of having been compiled by an experienced teacher of analysis. If another book of this sort be wanted for students preparing for the Science and Art Department, Medical Schools, London University, and Oxford and Cambridge Local Examinations, Mr. Eltoft's work may prove useful: at least, it may be said to present

some commendable features. But the effort after brevity and conciseness leads here, as elsewhere, to imperfections if not to errors. For instance, we do not see that any provision is made (pp. 25, 30, 53, and 55) for testing for fluorine when silicon is also present; the mode given for identifying oxalic acid is not conclusive; and the directions for removing, by means of sodium carbonate, heavy metals previous to testing for acids, are vague. A tabular statement of the general scheme of analysis for bases is omitted, although the substance of such a scheme is given in a less advantageous shape. Mr. Eltoft claims originality for the table of processes to be used in detecting acids (page 53), but we fail to see novelty in it. The Table of Solubilities, on page 9, does not strike us as very clear—indeed the 2nd section of it appears misleading.

Elements of Modern Chemistry. By A. WURTZ. Translated and Edited by W. H. GREENE, M.D. Pp. 687. London and Philadelphia, 1879.

STILL another manual of chemistry for students! We confess to having experienced some degree of disappointment on perusing this recent addition to the crowd of similar digests of chemical facts and theories. The great reputation of M. Wurtz, and the popularity of the original work in France, led us to expect a volume perfect as far as it went in its scope and in its selection and treatment of details. No doubt the philosophical or theoretical aspect of chemical science is fairly handled in these pages, but of the descriptive part of the work we find it impossible to speak in terms of equal praise. We object to the non-metals being called (p. 3) "metalloids"; we object to the statement (p. 7) that the objects around us present "infinite differences": when we look up particular elements and compounds, one after another, in order to test the lateness of the information given and the judgment shown in its selection and arrangement, we find old data and statements cited as if they were still of real value; we fail to discover that perfection of orderly arrangement which a systematic treatise, however compact, should always exhibit; and we here oftentimes search, in vain, for a notice of the most salient features and most characteristic data of the bodies described. One would imagine, for instance, that the specific gravity and hardness of such an important substance as rock-crystal would be given (p. 199–200), and that we should be spared such an inane remark as this, "chalcedony, onyx, and opal are sought for by the lapidary and engraver." One would expect, too, that more recent information about graphite would be furnished to us than the imperfect, half-obsolete, and partially incorrect statements of pages 201 and 202; surely, the existence of graphitic acid might have been noted. Reaching the metals, we do not fare much better. Under potassium (p. 282) the explanation offered to account for the explosive violence with which, when the metal burns on water, the resulting fused potassium hydrate is suddenly quenched and dissolved, includes no reference to the spheroidal state. Sodium hydrate, we are informed (p. 292), "is known in commerce as concentrated lye." Bleaching powder, according to the equation on p. 309, is made with anhydrous lime: we would ask how much chloride of lime is now manufactured in the manner described and figured (fig. 101) by M. Wurtz? On page 302, the author assigns to Lamy the honour, not only of establishing the metallic nature of thallium, but of first isolating that element. It seems to us impossible that any unprejudiced and well-informed chemist could honestly deny to Crookes the credit of having discovered thallium, and also of having first made known its metallic nature and first publicly exhibited it. Let any one who doubts this conclusion read the *Philosophical Magazine* for July, 1863. He will then find that Mr. Crookes exhibited thallium in the International Exhibition of 1862, on May 1st, duly labelling his case with the words "Thallium, a new metallic element, discovered by means of spectrum analysis." The individual specimens of metallic thallium and its salts were

there shown with labels which were proved to have been printed on the 25th of April, 1862. And what publication of Crookes's discovery could have been more conspicuous and incontrovertible than the exhibition of a series of duly labelled specimens to an immense concourse of visitors on an occasion like that of the opening of the Exhibition on May 1st, 1862?

We have not space to discuss the remainder of this book; the organic section especially would require more attention than we can give it here. But it appears from casual dips which we have taken into the latter part of this "Introduction to Modern Chemistry," that the new carbon compounds therein described are treated more satisfactorily than the older and more familiar ones. For instance, the description of malting and the figure of a sprouted barley grain, given on page 579, hardly commend themselves to the critical faculty. Cellulose when pure has a definite specific gravity, not one oscillating (p. 585) between 1.25 and 1.45; *Charpie*, too (pp. 585 and 586), has an English equivalent; and we were not aware that collodion was a solution of trinitrocellulose. Considering the paramount importance of chlorophyll in organic nature, we might at least expect one line about its occurrence and function. But, in fact, the details of this work do not bear the kind of scrutiny which one has a right to apply to a book written by Wurtz.

CORRESPONDENCE.

NEW ELEMENTS.

To the Editor of the Chemical News.

SIR,—The number of "new elements" announced in successive numbers of the CHEMICAL NEWS is really appalling. During the past two years no less than eleven claimants for recognition as elementary substances have presented themselves. With your permission we will call the roll, and inspect briefly the chemical recruits.

List of New Elements and their Discoverers.

1877, June	Davyum	Sergius Kern.
" "	Neptunium	Hermann.
" "	Lavoesium	Prat.
" May	Mosandrum	J. L. Smith.
1878, Sept.	"New earths"	Gerland.
" Oct.	Philippium	Delafontaine.
" Nov.	Ytterbium	Marignac.
" Nov.	Decipium	Delafontaine.
1879, March	Scandium	Nilson.
" July	Norwegium	Dahl.
" Aug.	Uralium	Guyard.

Now, while several names in this formidable list are doubtless destined to stand the test of time, the majority, I opine, will have to join the noble army of "defunct elements." Probably only the "fittest" will "survive."

Seriously, however, have we not a right to hesitate before accepting these claimants and granting them a place in our catalogue of elements. To put a practical question, How long shall we delay before mentioning these substances in the instruction of advanced classes? Shall we enter them in our text-books? Who will undertake to say that one chemist is thoroughly reliable, and that another is economical of truth or easily deceived? What authority is weighty enough to settle the existence or non-existence of these bodies?

These and similar thoughts doubtless occur to many. We would like to see the views of some of your correspondents on this knotty problem.—I am, &c.,

SCEPTICAL CHYMIST.

The Chemistry of Food.—A course of ten lectures on the Chemistry of Food will be delivered at the National Training School for Cookery, Exhibition-road, South Kensington, by Prof. Church. The lectures will be given on Mondays at 5 p.m., the course beginning on Monday next, the 15th instant.

MISCELLANEOUS.

Action of Pyrogallate of Potassa upon Binoxide of Nitrogen.—G. Lechartier.—The binoxide of nitrogen seems to undergo a decomposition, giving rise, on the one hand, to protoxide of nitrogen, and, on the other, to the more highly oxidised compounds, nitric and nitrous acids, whose action upon the pyrogallate of potassa is already known.—*Comptes Rendus*.

Synchronism of the Mean Temperature and Rain-fall in the Climate of London.—Mr. H. Courtenay Fox, M.R.C.S., of Stoke Newington, read a paper on this subject before the British Association at the Sheffield meeting. The paper was accompanied by copious tables exhibiting the months and seasons for sixty-seven years, arranged in the order of their respective temperature and rainfall as for the Royal Observatory. The following is a short summary of the results:—1. In each of the four months from November to February, extreme cold tends to be synchronous with dryness, warmth with large rainfall. 2. In the summer months, June to August, cold tends to be accompanied by much rain, warmth by dryness. 3. To put this in popular language, rain brings warmth in winter and cold in summer; that is (if rain be the cause, which is by no means proven) it mitigates the special character of each extreme season, winter and summer. 4. Very wet years tend to be either cold or warm, whilst years of drought tend to assume an average temperature.

TO CORRESPONDENTS.

W. S. S. G.—There is no process by which the odour can be removed from nitrobenzol.

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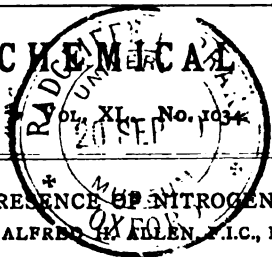
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THE CHEMICAL NEWS.



ON THE PRESENCE OF NITROGEN IN STEEL.*

By ALFRED W. ALLEN, F.I.C., F.C.S.

THE author made some preliminary experiments on this subject in 1872, but has only recently obtained any definite results. The method adopted has been to dissolve the steel in hydrochloric acid, by which means any combined nitrogen may be presumed to be converted into ammonia. The solution obtained was then distilled with excess of lime, and the distillate examined for ammonia by Nessler's method. The employment of this extremely delicate test enabled the author to operate on a much smaller quantity of steel than was employed by previous investigators. Very special precautions were taken to obtain the hydrochloric acid and other materials free from any trace of ammonia or nitrous compounds, and the air was entirely expelled from the apparatus before commencing the operation. The hydrogen evolved was freed from any traces of ammonia by passing it through a tube filled with glass beads moistened with hydrochloric acid. It was proved by blank experiments that no source of ammonia existed in the reagents or apparatus.

When absolutely pure materials were used, and every precaution taken to get rid of the contained air and other sources of error, the addition of Nessler's solution to the liquid obtained on distilling with lime caused a very marked yellowish brown colouration. On comparing the tint produced with that yielded by a dilute solution of ammonium chloride of known strength, results were arrived at representing the proportions of nitrogen present in various typical specimens of steel.

As the results obtained from steels of different kinds varied greatly, it cannot be assumed that there was a constant source of error in the mode of manipulation; while as the same samples gave substantially concordant results on repeating the experiment, the figures obtained are not the result of accident, but are true expressions of the proportions of nitrogen present.

In order to obtain ammonia in quantity sufficient for its recognition by other reactions than that with Nessler's test, the following plan was employed.

Steam, generated by boiling water in a flask, was passed over a considerable quantity of steel borings contained in a combustion-tube which was bent beyond the furnace, and prolonged so as to form the inner tube of a Liebig's condenser. To the further end, a tube filled with glass beads and furnished with a glass stop-cock was attached. A rapid current of steam was driven through the apparatus for a considerable time to expel every trace of air. On condensing the steam, it was found free from any trace of ammonia. The steel borings were then heated to redness by a combustion furnace, and a rapid current of water passed through the condenser. The condensed steam, when tested by Nessler's solution, was found to contain abundance of ammonia, which did not diminish in amount till the borings were almost entirely oxidised. On re-distilling the condensed steam, a distillate was obtained, having a distinctly alkaline reaction to litmus paper, and on treating it with hydrochloric acid and platinic chloride a sensible amount of yellow precipitate was obtained, having the characteristic crystalline form of ammonium chloroplatinate. The amount found was larger than could possibly have been produced had the whole of the nitrogen of any residual trace of air been converted into ammonia.

* Abstract of a paper read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

The author regards the results now recorded as preliminary merely, and proposes to extend the research to various classes of steel and iron, and especially to such specimens as have been found to possess anomalous characters. Of these characters, the evolution of ammonia from freshly fractured surfaces is among the most striking.

A HISTORICAL SKETCH OF VAPOUR-DENSITY METHODS.*

By JAS. T. BROWN, F.C.S.

ALTHOUGH Southern in 1803 made some very careful determinations on steam, still the first accurate method was that of Gay-Lussac, who, in 1811, heated a weighed quantity of substance over mercury in a graduated vessel. In 1822, Cagniard de la Tour studied the combined effect of heat and pressure on various liquids, but Despretz, in the same year, worked on vapours in exhausted globes of 9 litres capacity. In 1826, Dumas, wishing to operate on substances which attack mercury, worked out and published his well-known method, in which the *volume* is definite but the amount of substance used has to be determined. In 1833, Mitscherlich proposed using tubes, sealed at one end and drawn to a neck at the other, instead of bulbs, and gave details and drawings of the apparatus for heating them. Bineau, in 1838, published an elaborate paper, but unfortunately without any drawings; in 1846 he repeated the experiments of Despretz with slight modifications. In 1849, Regnault used an apparatus very similar to that of Mitscherlich, but closed the air-thermometer tube by means of a stop-cock. Bineau, in 1859, for high temperatures, used glass tubes coated with clay; but Regnault, in 1861, to obtain the same result, used iron tubes, and heated them in a revolving cast-iron tube over gas-burners. Another method of Regnault's was to employ cast-iron bottles with loose stoppers, and to use the vapour of mercury as the standard instead of air. In 1866, Grabowski used the original Dumas method, but for air-thermometer employed a globe of the same size as that containing the substance. Bunsen, in 1867, employed an air-bath similar to that of Mitscherlich, but heated it by an elaborate arrangement of gas-burners. For high temperatures Deville and Troost, in 1860, heated the bulbs in the vapour of mercury, sulphur, zinc, or cadmium; above the boiling-point of sulphur they used porcelain globes. For temperatures up to that point the smaller apparatus devised by Greville Williams answers admirably. Roscoe, last year, used porcelain globes and heated them in a muffle, but determined the temperature by the method of specific heat. For working at a reduced pressure Regnault proposed partially exhausting the bulb during the experiment; in 1876, Habermann (whose method was referred to by Sommaruga) gave a diagram of the complete apparatus. The main point in Playfair and Wanklyn's method (1861) of vapours mixed with air, consisted in stopping the supply of vapour before the bath had attained its maximum temperature.

Natanson, in 1855, to work up to 300°, heated the upper part of the Gay-Lussac tube in an air-bath by means of charcoal; he used Avogadro's tables for the tension of mercury vapour. In Greville Williams's (1857) apparatus for working at various pressures, the metallic cistern, by which communication was made between the bottom of the graduated tube and the bottom of the open tube, was small, so that the whole arrangement could be placed in the water- or oil-bath. In Regnault's apparatus for the same purpose the two tubes are fastened to the bottom of the water-bath, and are connected underneath by a T-piece. For ordinary determinations up to 150°, Greville Williams proposed standing the graduated tube in a small iron cup

* Abstract of a paper read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

containing mercury, and supporting it in a glass vessel resembling a large test-tube. Schiff, in 1862, proposed fastening a loaded handle to the upper extremity of the graduated tube by spring clips. Grabowski, in 1866, heated the tube in an air-bath by means of gas, and placed an exactly similar tube containing air by the side of the substance-tube, so that the comparison could be made direct between the vapour and air under the same conditions. Croullebois, in 1874, proposed using a tube with a large globe at the end, but Deville criticised his method rather severely.

In 1868, Hofmann adopted a tube more than 760 m.m. in length, and enclosed it in a slightly larger mantle tube. He then passed the vapour of a liquid of definite boiling-point through the intervening space, selecting the liquid according to the temperature required. He thus maintained a steady temperature with the greatest ease. Wichelhaus, in 1870, fitted a modified U-tube to the bottom of the graduated tube, so that the whole of it could be surrounded by vapour. Grabowski, in 1875, used naphthalen with Hofmann's apparatus, but Engler, in the following year, to avoid the occasional stoppage of the outlet-tube, fitted to the lower end of the outer tube a short side tube, similar to those used for heating funnels. He then boiled the heating medium in this tube and allowed the vapour to cohabit in the space between the two glass tubes. Hofmann, at the same time, proposed making the outer tube long enough to enter the mercury in the trough, and placed the outlet a short distance above the mercury-level. He also proposed using plain tubes instead of graduated, by marking the height of the column by a gummed label and then measuring the volume occupied by the vapour. His third proposal was to place the inlet and outlet both at the bottom of the outer tube, so that during the heating of the liquid in the boiler, a portion of it could be forced at will into the space between the two glass tubes so as to heat them more gradually.

Brühl used an inner tube 1.5 m. in length, and to eliminate the tension of mercury vapour heated the column to the required temperature, noted the height, allowed it to cool, introduced the substance, and then heated again to the same temperature. He also made a mark on the tube a little above the vacuum mercury-level, and then only calibrated about 150 m.m. down from that point. Muir and Suguira, in 1877, ensured communication between the mercury in the tube and that in the trough by means of a short piece of glass tubing bent at right angles, which passed through the india-rubber disc on which the tubes rested. A second tube long enough to stand slightly above the level of the mercury in the trough served as outlet from the space between the two tubes. Brühl has this year proved by most carefully conducted experiments that the Hofmann method cannot be used above 220°, owing to the great and rapidly increasing vapour-tension of mercury. Of the overflow methods, the first is that of Hofmann, who, in 1860, wrote that he used an U-tube heated in a paraffin bath, and estimated the volume of the vapour by the mercury expelled. Wertheim (1862-4) gave full details of his method, in which he used two short tubes suspended in a flask. Watts, in 1867, employed a globe with a ground neck into which an outlet tube reaching nearly to the bottom of the globe was accurately fitted. Victor Meyer, in 1876, used fusible metal and placed the outlet at the bottom of the bulb. His experiments were all made in the vapour of boiling sulphur, but Graebe last year used phosphorus pentasulphide. Friedrichs, in 1876, used an inverted flask and brought the outlet through the bottom of the bath. Goldschmidt and Ciamician, in 1877, used mercury with the simpler bulb of V. Meyer, but added a small side tube to the outlet, so that the mercury expelled could be weighed during the heating. V. Meyer, in the same year, modified the shape of the bulb, but heated it in a tube of sufficient length for the upper part to serve as condenser. Pfaunder's method (1870 and 1879) is based on the increased tension of the air, in an elongated bulb, produced by heating after the

introduction of the substance, as compared with a similar determination on air in a bulb of the same size. A very short description appeared in 1874 of a method by Dulong on the same principle. Last year Hofmann proposed (1) to heat the weighed substance over mercury in the closed limb of an U-tube, and to calculate the volume from the extent to which heating caused the mercury to rise in the open limb; (2) to heat a limited but weighed quantity of substance in an exhausted tube, and to calculate the volume of the vapour from the amount of air which entered the tube to restore equilibrium while at the maximum temperature. V. and C. Meyer's method, which is so recent and so well known as not to require any explanation, is based on the measurement under atmospheric conditions of the air expelled by the vapour, the substance being introduced *after* the bulb has been heated.

THE PHYSICAL PROPERTIES OF LIQUID ACETYLEN.*

By GERRARD ANSDELL, F.C.S.,
Chemical Assistant to the Royal Institution of Great Britain.

THE hydrocarbon acetylen, being the only one of its numerous class which can be formed synthetically by the direct union of its constituent elements, has an especial interest for the chemist. The chief physical constants of this substance are unknown, although its polymerised modification, benzene, has been very thoroughly studied in its physical relations by Regnault. Having one of M. Caillietet's ingenious pumps for the liquefaction of gases at my disposal in the laboratory of the Royal Institution, Professor Dewar suggested a series of accurate determinations of its physical properties in the liquid state, and the present communication deals with the critical point, the tension of the vapour of the fluid at various temperatures, together with the corresponding densities and coefficients of compressibility.

The only notice on the liquefaction of the gas appears to be a short paper by M. Caillietet in the *Comptes Rendus*, vol. lxxxv., No. 19, in which he determines the tension of the vapour of the liquid at different temperatures. These tensions, as will be afterwards seen, differ entirely from those obtained in the present paper, one of the reasons appearing to be that instead of using a carefully calibrated air manometer for determining the pressures, he used the ordinary metallic gauge attached to the pump, which is far from being correct.

The pump itself is too well known to need description; suffice it to say that two of the iron bottles or reservoirs were used, connected with the pump by a piece of fine-bore copper tubing, so as to equalise the pressure, one containing an air manometer registering the pressures from ten atmospheres upwards, and the other the tube filled with acetylen. The two bottles were then placed by side, and the height of the column of mercury in either read off by means of a cathetometer.

The formulæ used for calibrating the tubes, and also for calculating the volume of the liquefied gas, and the pressure by the air manometer, were those given by Dr. Andrews in his researches on carbonic acid (*Phil. Trans.*, 1869 and 1876). The method of preparing the acetylen gas was by the action of alcoholic potash on bibromethylen, the disengaged gas being collected in the form of the red acetylide of copper, by passing it into a strong solution of subchloride of copper in ammonia. This red compound, after being thoroughly washed and boiled with distilled water, was transferred to a flask with dilute hydrochloric acid, the gas driven off by means of a gentle heat, and conducted through a strong solution of caustic soda, to free it from traces of hydrochloric acid, and finally

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

through two small U-tubes with fused chloride of calcium. The perfectly pure and dry acetylen was now passed through the tube to be used for its liquefaction in a slow stream for several hours, and the latter carefully sealed off when all the air had been expelled.

The sealing off requires great care, as unless rapidly done, and the pressure removed from the inside (by cooling the tube) immediately the point is closed, the acetylen becomes rapidly charred and blows out, a small portion of it being consequently decomposed, and thus interfering materially with the accuracy of the results.

The tube for the tension determinations was of the usual shape used in the Cailletet pump, the internal diameter of the capillary part being about 2.5 m.m. This was found to be more convenient than a narrower tube, as a larger reservoir could be used, and consequently a larger quantity of liquid obtained.

The pressure at the different temperatures was always observed when a very slight layer of liquid was formed on the surface of the mercury, as the gas not being entirely free from air, the pressure was slightly increased on filling completely the upper part of the tube.

The following are the tensions obtained compared with those of Cailletet:—

Cailletet.			
Temp.	Pressure.	Temp.	Pressure.
-23.00° C.	11.01 atm.		
-10.00	17.06 "		
0.00	21.53 "	+ 1.0° C.	48 atm.
+ 5.25	25.48 "	2.5	50 "
13.50	32.77 "	10.0	63 "
20.15	39.76 "	18.0	83 "
27.55	48.99 "	25.0	94 "
31.60	56.20 "	31.0	103 "
36.00	65.36 "		
36.50	65.89 "		
36.90	67.96 "		

The temperatures above zero were kept constant to within 1/10th of a degree by allowing a constant stream of water to flow over the tube from a reservoir holding about 10 gallons, in which it had been previously thoroughly mixed. The temperature of -10° was obtained by cooling down alcohol with ice and salt, and that at -23° by surrounding the tube with a narrow glass cylinder containing liquid chloride of methyl, which boils constantly at this temperature; this cylinder being again enclosed in a wider one containing a little phosphoric anhydride to prevent moisture from condensing on the sides.

It was thought interesting to compare the tensions of liquid acetylen with those of the saturated vapour of benzene, being polymeric bodies, although having totally different properties. For this purpose curves were plotted for the two substances, that for the benzene being taken from Regnault's results (*Mém. Acad. Sci., Paris*, vol. xxvi., p. 420). They do not, however, run parallel to each other, the benzene having a slower rate of increase at low temperatures, but a quicker rate than the acetylen as the temperature rises. The curves, however, have no appearance of actually crossing at a higher temperature.

The critical point of acetylen, or that temperature at which no appearance of liquefaction takes place, however great a pressure is exerted on the gas, was found after many careful experiments to be 37.05° C.

For determining the density and compressibility of the liquid at different temperatures a tube of much smaller dimensions was used, having a capillary bore of about 0.8 m.m. in diameter, the whole of the tube having a capacity of 36.3708 c.c. This gave a column of liquid about 15 centims. long when the upper part of the tube was entirely full at 15° C.

The density at any particular temperature was taken by forcing the liquid up the capillary tube at that temperature until the upper part was completely filled. The length of the column of liquid was then read off, its volume calcu-

lated, and this observed volume divided into the calculated weight of the gas at zero. They are as follows:—

Temp.	Density:
- 7.00° C.	0.460
- 3.00	0.456
0.00	0.451
+ 4.40	0.441
9.00	0.432
16.40	0.420
20.60	0.413
26.25	0.404
30.00	0.397
34.00	0.381
35.80	0.364

It has, therefore, about half the density of liquid carbonic acid, and if we take the actual volume of the liquid at -7 as unity, it becomes 1.264 at +35.8, which gives 0.00489 as its coefficient of expansion per degree for the total range of pressure; it is, therefore, only about half as expansible as carbonic acid, whose coefficient is 0.010, and is not much more expansible than a gas. Comparing the density of liquid acetylen with that of liquid benzene, the latter is found to be almost exactly three times as great as the former at the same temperature; as, for instance, at 0° C. the density of the acetylen is 0.456, whereas that of the benzene is 0.899. The vapour-density, however, of the benzene is three times as great, viz., 2.704.

The apparent compressibility in glass was determined by direct observation, the liquid being forced up in the capillary tube until the latter was completely full, and then the pressure gradually increased, and the diminution of volume read off at intervals of about 10 atmospheres up to about 180 atmospheres.

Curves were then plotted, showing the volume at different pressures for the same temperature, and from these the coefficient of compression at any temperature and pressure was easily deduced.

The following tables are constructed from the curves:—

I. Mean Coefficients of Compression of Liquid Acetylen at Different Temperatures. Range of Pressure from 36.62 to 182.68 atmospheres.

Temp. of Acetylen.	Coeff.	Temp. of Acetylen.	Coeff.
35.0° C. =	0.00085	16.0° C. =	0.00050
28.6 =	0.00068	4.4 =	0.00038
22.5 =	0.00058	0.0 =	0.00025

II. Coefficients of Compression at the same Pressure but varying Temperatures.

Temp. of Acetylen.	Atm. 90.	Atm. 95.	Atm. 120.	Atm. 160.
*49.0° C.	—	0.00343	0.00169	0.00078
*41.0	—	0.00138	0.00099	0.00076
35.0	0.00171	0.00113	0.00078	0.00065
28.6	0.00122	0.00083	0.00072	0.00050
22.5	0.00079	0.00065	0.00057	0.00047
16.0	0.00066	0.00050	0.00049	0.00035
4.4	0.00047	0.00042	0.00034	0.00032
0.0	0.00041	0.00036	0.00025	0.00029

III. Coefficients of Compression at Varying Pressures and Temperatures Corresponding to the same Volume.

Temp. of Acetylen.	Vol. = 97 c.m.m.		Vol. = 92 c.m.m.	
	Pressure.	Coeff.	Pressure.	Coeff.
*49.0° C.	170.8 atm.	0.00080		
*41.0	137.0 "	0.00085		
35.0	103.2 "	0.00093	175.8 atm.	0.00065
28.6	70.0 "	0.00120	137.8 "	0.00063
22.5	—	—	99.2 "	0.00065
16.0	—	—	59.5 "	0.00066

* These two experiments were, of course, made above the critical point.

	Vol. = 101 c.m.m.	Vol. = 89 c.m.m.
*49°0	126°3 atm. 0°00128	
*41°0	98°3 " 0°00132	
35°0	72°7 " 0°00167	
22°5	— — —	158°0 atm. 0°00054
16°0	— — —	115°6 " 0°00056
4°4	— — —	49°7 " 0°00058

It is evident from the above tables that acetylen is governed by the same laws as other compressible liquids; that is to say, its compressibility increases as the temperature rises, but diminishes as the pressure increases. For instance, at a pressure of 95 atmospheres it is three times as compressible at 35° C. as at 0° C.

The volume being the same, the compressibility appears to be nearly the same at different temperatures, which is really due to the curves at high pressures running nearly parallel, thus introducing a corresponding difficulty in the estimation of small differences.

On comparing the compressibility of liquid acetylen with the results obtained by M. Amagat (*Annales de Chem.*, 1877) in the case of benzene, it appears to be about seven times as compressible as the latter body at a temperature of 16° C., and under a pressure of 40 atmospheres. The comparison could not be carried out at higher temperatures, for whereas M. Amagat reaches a temperature of 100° C. with the benzene, I was not able to go beyond 35° C. with the acetylen.

NOTES ON A SAMPLE OF FULLER'S EARTH FOUND IN A FULLONICA RECENTLY EXCAVATED AT POMPEII.*

By WILLIAM THOMSON, F.R.S.E.

In visiting the ancient city of Pompeii in April last, I observed in one of the Fullonica establishments a large square tank set in the ground filled with a white soft substance which was soapy to the touch, and which was pointed out to us as the soap of the ancients. I took a sample of the substance with a view of making a chemical examination of it.

This substance is named by the Italians "Terra Fullonica," and besides being found in the dyers' and washers' quarter of the city, it has been discovered frequently in the ordinary houses which have been excavated.

Among the literature of chemistry I searched for, but failed to find, any mention of this fuller's earth or of its composition, but through the kindness of Sig. Felice Niccoline, Director of the National Museum of Naples, I obtained a pamphlet written by Professor de Luca, entitled "Chemical Researches on 'Terra Fullonica,' found in Pompeii, April 13, 1878."

In this he gives general chemical peculiarities of the clay, such as its being faintly alkaline and containing silica, lime, magnesia, chlorides, and traces of sulphates—potassium and sodium. He gives its composition according to a mechanical analysis as follows:—

50 grms. of the clay stirred in water gave different residues, which were separated according to their tendency to settle to the bottom.

10°282 grms. settled first, and was composed of sand and carbonate of lime.

17°710 grms. was composed of a little sand and carbonate of lime and much clay.

10°050 grms. is formed of traces of carbonate of lime and much clay; and the fourth residue,

8°230 grms., was greasy to the feel, and plastic, and fused before the blowpipe into a vitreous bead of a yellowish white colour.

Prof. de Luca also states that it contains 17 per cent of water, 24 per cent of matter soluble in hydrochloric acid, and 2·7 per cent carbonic acid, the remainder being insoluble in hydrochloric acid.

On drying the substance thoroughly at 100° C., and then submitting it to analysis, I found it to be composed of—

	Per cent.
Silica	67°145
Alumina	12°857
Oxide of iron	2°107
Lime	6°412
Magnesia	1°822
Carbonic acid	3°451
Manganese	trace
Combined water	3°953
Alkaline salts and loss	2°253
	100°000

PHOSPHORUS IN ANCIENT IRON.

By A. E. ARNOLD, A.I.C.

In former times it has been observed, when iron was made in small forges and at comparatively low temperatures, in a similar manner to that still employed in certain half civilised countries, the greater part of the phosphoric acid passed into the ferriferous slag, and good malleable iron was finally produced tolerably free from phosphorus. The following is an analysis of a scoria produced in Roman or Etruscan times from the specular ore of Elba, which usually contains about 0·04 per cent of phosphoric acid.

Iron protoxide	76°49 per cent
Alumina	2°57 "
Manganese protoxide.. .. .	0°60 "
Lime	1°32 "
Magnesia	0°64 "
Zinc oxide	0°20 "
Sulphur	0°40 "
Phosphoric acid	0°34 "
Silica	14°20 "
Oxygen and not estimated.. .. .	3°88 "
	100°00 "

This specimen was taken from a large heap of many thousands of tons of scoria lying on the beach close under the site of Populonia, the Etruscan Pupluna, a town much famed in those times for its iron and copper manufactures. It furnished Scipio with iron in the second Punic war. It is now a deserted site, and but few traces of its former importance remain.

It is of interest to observe in the above analysis that the phosphoric acid is about eight times as much as in the natural ore.

I obtained in February, 1878, a piece of metallic iron from the same district; it originally weighed about 2 kilos., and was somewhat rusted. It was found among scoria in the vicinity of Campiglia, and gave on analysis:—

Combined carbon	0°873 per cent
Graphite	2°853 "
Silicon.. .. .	0°544 "
Sulphur	0°096 "
Phosphorus.. .. .	0°090 "
Manganese.. .. .	0°091 "
Iron sesquioxide	2°430 "
Metallic iron	92°804 "
Moisture	0°092 "
	99°873 "

Of the antiquity of this specimen there is some doubt but it has certainly not been produced in recent times.

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

The high percentage of carbon cannot be adduced as indicative of its modern origin; the researches of Lowthian Bell proving that carbide of iron in contact with iron oxide in the molten state is not necessarily decarburised. Similarly, therefore, the above specimen might have been made contemporaneously with a richly ferruginous scoria, such as the Roman or Etruscan metallurgists produced. The ancients possibly knew iron both cast and wrought, as well as the intermediate steel. Pliny says that iron was made in a similar manner to copper. Making a comparison between puddling and copper smelting, he notes the "remarkable fact that when the ore is fused the metal becomes liquefied like water, and afterwards acquires a spongy brittle texture" (xxxiv., 41), which goes to imply that iron first "came to nature" considerably previous to the days of Mr. Cort.

Cast-iron recently produced from Elba ore near the same locality contained three times the quantity of silicon present in the above analysis, the constituents being as follows:—

	I. Per cent.	II. Per cent.	Mean. Per cent.
Carbon	4.306	4.147 (diff.)	4.306
Silicon	1.672	1.676	1.674
Sulphur	0.067	0.056	0.067
Phosphorus ..	0.110	0.108	0.109
Manganese ..	0.748	0.757	0.753
Iron	93.256	93.256	93.256
	100.159	100.000	100.165

8, effrey's Square, London,
September 10, 1879.

THE AMMONIACAL COPPER-TEST AND ITS APPLICATION.

By J. STEINER.

My attention having been drawn to the working of the ammoniacal copper test (as recommended by Dr. Pavy) by Mr. Hehner's remarks, I determined to find out whether this process could be applied with advantage to the estimation of sugar in malt liquors, especially when old, black, or "turning" beers are to be analysed. The volumetric determination, as carried out in the usual way with ordinary Fehling's solution, does not give in these instances quite satisfactory results in consequence of the appearance of a deep straw-yellow colour after boiling for some time and when about three-fourths of the total cupric oxide have been reduced. Moreover, the recourse we have to testing a few drops of the filtered and neutralised solution with ferrocyanide fails here because much ammonia is developed from the albuminous matter present, and therefore part of the Cu_2O remains in solution. Just the same is the case when the amount of glucose or "uncrystallisable sugar" in Jaggery, China or Jamaica sugars has to be ascertained, and in other instances, —especially with old pale ale or old lager beer, where the sugar in the beer is reduced by after-fermentation to less than one-half per cent.—unless the beer solution be added gradually to the copper test, and the boiling is repeated several times, a muddy hydrated Cu_2O is separated, and remains suspended in the liquor, in which case even filtering is useless, and the test has to be put aside. In carrying out my experiments with an ammoniacal copper solution as prepared by Dr. Pavy or Mr. Hehner, I soon perceived that the different measurements for the two tests impeded the close comparison of the results; and as for the usual sugar test I apply 10+10 c.c. = 20 c.c. of a copper solution, the parts of which I keep in two separate bottles; I repeated the test with the same amount of test-liquor after the addition of about 30 to 40 c.c. of strong ammonia, and with or without 10 c.c. (= 1 gr.) of Na_2O . In this manner I arrived at very easily comparable results.

For the ordinary Fehling's test I use a test-tube of about 120 c.c. capacity. It is held direct over a Bunsen flame, and the bumping of the dense liquor is prevented by the addition of a few grains of pumice stone. This very convenient mode of procedure I saw first in Mr. Halse's laboratory, in London. The ammoniacal test I carried out in a boiling flask of about 200 c.c. capacity. It was closed with a perforated cork, and the fumes carried into cold water. It is very essential to prevent an access of air, as the reduced colourless solution very soon becomes blue on lifting the stopper. The beer or the sugar solution was so diluted as to require about 35 to 45 c.c. of the same for 20 c.c. of Fehling. I used, first, inverted pale cane-sugar, and came very soon to the conclusion that 20 c.c. of the copper test corresponded to—

- 1-10th gr. glucose, in the ordinary mode of testing.
- 1-12th gr. glucose, with the addition of ammonia+1 gr. of Na_2O .
- 1-12.5th gr. glucose, with the addition of ammonia only.

For example:—Required for the ordinary process = 36 c.c. (I.) of sugar solution, but with the addition of ammonia+1 gr. Na_2O =30 c.c. (II.), and with ammonia only=28.8 c.c. (III.).

It will be noticed that these numbers stand to each other in the reversed proportion of 10:12:12.5; and that 100 c.c. of this sugar solution contain according to—

- (I.) 36:0.1=100:($x=0.2778$) gr. glucose.
- (II.) 30:1.12=100:($x=0.2778$) " and
- (III.) 28.8:1.125=100:($x=0.2778$) "

The same proportion was obtained with inverted brown cane-sugar and with brown starch sugar (English make).

It is worth noticing here that while the ordinary test requires, in presence of dextrin, a rapid working and not a too protracted boiling (or the results become too high), the ammoniacal test, with or without caustic soda, is best carried out gradually with a low flame, to prevent the adhering and decomposition of the test-liquor at the bottom of the flask, and the results are not influenced whether boiling rapidly or only slowly is resorted to. I next used a strong malt wort from the mash in the brewery, diluted 20 c.c. of it to 1000 c.c., and to my surprise, in all the three instances, the same amount of c.c. (=40) was required. I then looked into Mr. Hehner's paper (CHEMICAL NEWS, vol. xxxix., p. 197), and noticed the following statement:—"Dextrose and levulose act equally, but I find that both milk-sugar and maltose possess entirely different reducing power against the two kinds of copper solutions. Thus, a solution of pure milk-sugar acted as follows:—10 c.c. ordinary Fehling required 16.76 c.c., 100 c.c. of Pavy solution =20.76 c.c. The ratio is therefore quite contrary to that resulting by the action of glucose."

There is, of course, a misconception here, because if it is considered that 100 c.c. of Mr. Hehner's ammoniacal copper solution equal 12 c.c. of ordinary Fehling, and the reduction from 12 to 10 c.c. in the proportion 12:10=20.76: x is carried out, we arrive at 17.3 c.c. against 16.76 c.c. in the usual way, with a difference of only 0.24 c.c., and which may be accounted for by the measurement of different quantities used by him. It follows, therefore, from his results that lactose is as little influenced as it is, according to my figures, the case with maltose when ammonia and excess of alkali are added to the common copper solution.

Having established these facts, I now tried a sample of American glucose, which was very white, and judging from the large quantity of the ash and its character, it was manufactured from maize. The result of the two tests, taken together, was that only dextrose was present in this sample; but a glucose of German manufacture, which was also very pale and contained only little ash, gave results which were not concordant. Of 1 gr. dissolved and made up to 250 c.c. were required, with ordinary Fehling's solution, = 36 c.c., and with the addition of

Am + 1 gr. $\text{Na}_2\text{O} = 32$ c.c., and on calculating the reduction as glucose we obtain—

In the one case, $36 : 0.1 = 250 : (x = 0.6944)$, or 69.44 per cent glucose.

And in the other, $32 : 1.12 = 250 : (x = 0.6510)$, or 65.10 per cent glucose.

This discord may also be noticed in the following proportion:— $12 : 10 = 36 : (x = 30$ c.c.), instead 32 c.c. as found by analysis, thus proving that there must be another sugar present, which in this instance is of course maltose.

The question now arose, How much maltose and how much glucose or dextrose is present? and to solve this problem I used the following equations:—

$$\text{I. } g + \frac{1}{2}m = F$$

$$\text{II. } 1.2g + \frac{1}{2}m = F'$$

From the combination of which follows—

$$g = \frac{F' - F}{0.2}$$

20 c.c. of copper solution are taken here for both tests to be equal to 0.1 gr. of glucose, and as the action of maltose (=m) remains the same for both cases, the difference in the obtained results (F and F') depends on glucose (=g), and the amount taken into consideration in the second equation (for the ammoniacal test) is to that (of the ordinary test) in 1. too high in the proportion of 1.2 : 1.0.

In applying this conclusion to the values given above, we have—

$$36 : 0.1 = 250 : (x = 0.6944) \text{ gr. and}$$

$$32 : 0.1 = 250 : (x = 0.7812) "$$

Therefore—

$$g = \frac{0.7812 - 0.6944}{0.2} = 0.434 \text{ gr., or } 43.4 \text{ p.c. glucose.}$$

From the equation—

$$m = \frac{3(F - g)}{2}$$

follows now—

$$m = \frac{3(0.6944 - 0.434)}{2} = 0.3906 \text{ gr., or } 39.1 \text{ p.c. maltose.}$$

The sample under examination contains, therefore, a total of 82.5 per cent fermentable sugars.

On inverting a solution of 1 gr. of this sample for six hours under pressure in a salt-bath I obtained, after diluting to 250 c.c., the following results with the two tests:—

I. 30 c.c. : 0.1 = 250 : (x = 0.8333), or 83.33 p.c. glucose.

II. 24.8 c.c. : 1 = 250 : (x = 0.8400), or 84.0 " "

Hence there was no dextrin present, and the sample consisted solely of glucose and maltose.

There is another mode of calculation, viz. :—

Let x be the number of c.c. of maltose and y be the number of c.c. of glucose which participate in reducing 20 c.c. of Fehling.

Then is $x + y = a$ c.c. for the common test,

And $x + \frac{10}{12}y = b$ c.c. for the ammoniacal and alkaline test,

Consequently $y = 6(a - b)$.

On applying this formula to the case considered above, we find $y = 6(36 - 32) = 24$ c.c., and as the total volume required in the ordinary test is 36 c.c., we learn from the proportion $36 \text{ c.c.} : 24 = 0.1 \text{ gr.} : (x = 0.0667)$ that out of the matter which reduces Fehling here 66.7 per cent is real glucose, and therefore the total amount of dextrose may be calculated from the following proportion:—

$$100 : 66.7 = 0.6944 : (x = 0.4631) \text{ gr., or } 46.31 \text{ per cent.}$$

For the maltose remains, therefore, $\frac{1}{2}(0.6944 - 0.4631) = 0.1157$ gr., or 34.7 per cent.

The two calculations did, therefore, not lead to the same result, and in order to obtain a better insight into this subject I mixed solutions of inverted cane-sugar with

diluted fresh malt-worts, and analysed several of these liquors before and after mixing them. The conclusion I arrived at now was that the fault lies, first, in the measurement, i.e., the burettes used for such work should permit an accurate reading off of $\frac{1}{100}$ c.c.; and, secondly, that the working of the ordinary Fehling's test requires to be most carefully watched, and repeated before the result may be noted down. In the subjoined data I give the details of such an experiment.

With Fehling. With Am. Copper Solution.

Dilute wort required . . . 35.6 c.c. and 35.6 cc.

Inverted cane-sugar required.. 46.3 " 38.6 "

50 c.c. of wort mixed with 50

c.c. sugar solution . . . 40.9 " 37.2 "

I. If we substitute the respective numbers in $y = 6(a - b)$ we obtain $y = 6(40.9 - 37.2) = 22.2$ c.c. for glucose, while the real amount is—

$$\frac{46.3}{2} = 23.15 \text{ c.c.}$$

II. 100 c.c. of the mixture contain, if expressed as glucose—

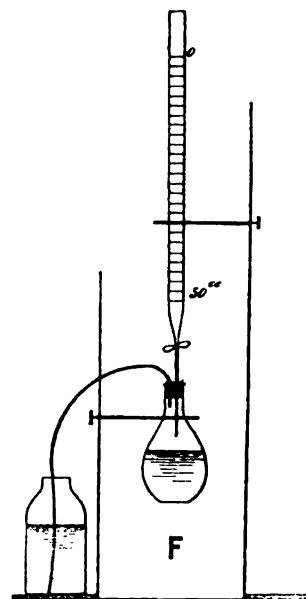
$40.9 : 0.1 = 100 : (x = 0.2445)$ gr. according to the usual copper test,

And $37.2 : 0.1 = 100 : (x = 0.2688)$ gr. according to the ammoniacal test.

Therefore—

$$g = \frac{0.2688 - 0.2445}{0.2} = 0.1215 \text{ gr.}$$

instead of 0.1080 gr. as the following proportion shows:—
 $46.3 : 0.1 = 50 : (x = 0.108 \text{ gr.})$.



The results obtained for such cases are therefore only approximate, and we have still to resort to the application of the polariscope, which, considering the large amount of substance used for a "normal solution," offers much more reliable data for calculation, and its indications ought never to be neglected in the analysis of starch-sugar. Nevertheless, I consider the ammoniacal alkaline copper test as most valuable for telling by merely analytical means whether a starch-sugar or a beer contains only one sugar or if two of them are present; and, as pointed out in the introduction of this paper, such analysis is very reliable if there be only one sugar, like in diabetic urine, or as in most beers and in raw sugars. In conclusion I will offer a few hints as to the *modus operandi*.

The apparatus used for this test is shown in the annexed sketch. The boiling flask should not be filled to more than two-thirds of its capacity. It is best first to ascertain with the usual test how many c.c. of the sugar solution are required, then to calculate from this according to the proportion 12 : 10 how much will be necessary for the reduction of the ammoniacal solution in the flask, and now to add at once before boiling the total amount less about 1 c.c. as a too protracted boiling drives out so much of the ammonia that Cu_2O is deposited and the test has to be repeated. The flame must be constantly in the operator's hand, and should not touch the bottom of the flask when the reduction once sets in, to prevent a burning and decomposition. The sugar solution must not be poured in while the flame is applied, or even while there are yet bubbles on the surface of the hot liquor in the flask, or the ammonia will pass up into the burette, alter the meniscus, and partly decompose the solution in it. The test is finished when a slight yellow tinge appears on looking on the surface of the hot liquor: a white cardboard as background aids the observation materially. The india-rubber tube is closed at the end, and a Bunsen valve, i.e., a longitudinal slit in it near the end, permits the gas to escape, and prevents the entrance of the cold water. This test has only one, but for many reasons a very objectionable, drawback, viz., the unavoidable penetrating smell of the ammonia. A water analyst, for example, could never make use of it in the same locality where the amount of ammonia in drinking-water has to be ascertained; or some of the analysts in the City will have to avoid it for fear of creating a public nuisance.

ON THE DISSOCIATION OF CHLORINE.

By F. P. DUNNINGTON, University of Virginia.

IN view of the following facts, the question suggests itself whether due care has been taken to exclude the possibility of oxygen compounds being present in the experiments, from which the decomposition of chlorine was inferred, in the article published in the *CHEMICAL NEWS*, vol. xl., p. 69.

During the past year in undertaking to make liquid Cl from PtCl_4 made by HNO_3 and HCl (the latter in excess) I experienced some difficulty in obtaining it free of nitro compounds.

After heating about 30 grms. PtCl_4 until Cl was given off, thrice it was treated with water, causing the evolution of nitrous fumes, and heated again until Cl was given off. This material sealed in a glass tube, on being further heated gave Cl, which condensed at first to a clear yellow liquid (exactly similar in colour to that of liquid Cl made from $\text{Cl}_2\text{H}_2\text{O}$), but on continuing the heat the condensed liquid assumed a red-brown tint. From the fact of the nitrous fumes having been evolved when it was last treated with water, I presume the reddish tint of this liquid due to NOCl or NOCl_2 . On opening the tube, when most of the Cl had escaped, there remained an odour very similar to that of ClO_2 .

In a previous experiment when the PtCl_4 was but once moistened after drying, the condensed liquid Cl had from the first this same tint, though to a more marked degree. In another experiment I did obtain the Cl from PtCl_4 wholly of the clear yellow colour.

In these experiments the residue after heating consisted of PtCl_2 , with but little Pt and PtCl_4 .

Detection of Phenol.—Dr. E. Hoffmann pours into a small test-glass 1 or 2 c.c. of pure concentrated sulphuric acid, pours carefully over it, so as to form a separate layer, the same volume of the dilute aqueous liquid suspected, and drops in a few granules of potassium nitrate. If only 1 milligram of phenol be present each particle produces at once violet streaks.—*Chemiker Zeitung*.

IRRIGATIONISM IN FRANCE.

It appears that certain engineers are making the attempt—if they have not already succeeded—to dispose of the sewage of Paris by converting the forest of St. Germain into an irrigation field. The question has been discussed before the Société d'Hygiène, and the debate is now continued in the *Journal d'Hygiène*, where M. Tollet and M. Duverdy have come forward in defence of the forest. They point out that it would be very unfortunate both for the salubrity and the amenity of the environs of Paris should this forest be cut down and rooted up. Although the Engineers of the city of Paris ask for one-third only of the forest, yet, as M. Duverdy very justly remarks, if once allowed to set foot in the forest they will speedily declare that one-third of the region does not suffice for the absorption of their sewage, and the remaining two-thirds will be in danger of the same fate.

M. Marié-Davy seems to believe that the trees might still be preserved even though the ground should be irrigated, and would even flourish more luxuriantly. This, however, is a capital error. The surface of the soil in a forest is not regularly turned over with the spade or the plough, and consequently, if drenched with sewage day by day, it will become felted up with the fibrous matters held in suspension, and be rendered impermeable. Very few forest trees can support a perpetual irrigation which must be applied all the year round, even during the season when the sap does not circulate. M. Marié-Davy, borrowing the expression of M. Durand-Claye, terms the part of the forest more immediately coveted by the engineers "an arid desert." M. Duverdy, on the contrary, shows that the trees, where not injured by rabbits, are healthy and luxuriant, and that the forest is valued by competent judges at 4200 francs the hectare,—a price which fully refutes the charges of aridity and barrenness.

In these days the sanitary value of trees in and near great cities is recognised by the highest authorities, and we trust the powers that be will not sanction an act of such wanton Vandalism. A palace burnt down may be re-erected, but a forest laid waste is almost irreparable.

NOTICES OF BOOKS.

Spon's Encyclopædia of the Industrial Arts, Manufactures, and Commercial Products. Edited by G. G. ANDRÉ, F.G.S. Division I. London: E. and F. N. Spon.

THE exact scope of this important work is not very easy to define. From the contents of this first division it might be taken for a dictionary of technological chemistry. The prospectus, however, promises to deal with many subjects not included among the chemical arts, such as hat, fan, and glove making, the manufacture of cotton, linen, silks, and woollens, &c. On the other hand, there is no mention of several important topics which could not well be overlooked in a work purporting to be a general encyclopædia of the useful arts. It seems to be the plan of the editor to discard what may be called the dictionary arrangement, and to group the information given under comparatively few heads, each of which will have to be carefully searched through by those in quest of any particular fact. The difficulty is enhanced by the circumstance that within these primary divisions the alphabetical arrangement is not always observed. Thus, in the chapter on acids we find the following compounds only described, and in the succession given: Acetic, arsenious, carbazotic (picric), carboic, carbonic, chromic, citric, gallic, sulphuric, hydrochloric, hydrofluoric, nitric, oxalic, and tartaric. It will at once strike the reader not merely that many important acids are omitted, but that the sulphuric acid is curiously out of place.

The salts are also distributed, some being placed under their respective acids, whilst others are appended to their basic constituents, *e.g.*, under the head "Alkalies." Hence we think that, especially as cross-references are not very abundant, a good index will be found absolutely necessary, and till it appears the value of the work cannot be fully appreciated.

As for the subject-matter of this encyclopædia we are happy to pronounce it very much superior to the arrangement, which, however, renders the latter more to be regretted. Many of the descriptions of manufacturing processes are accurately and carefully compiled, and are accompanied with abundance of illustrations. Still we cannot, in every case, agree with the writers. Thus, in the account of oxalic acid a process is described at very considerable length for its manufacture from guano, *i.e.*, from uric acid. We always regarded this invention as one of the curiosities of patent literature, unmatched save, perhaps, by a certain process for using alum as a source for potash and ammonia (Specification No. 938, A.D. 1855). In civilised countries uric acid is likely to be at all times much more costly than oxalic acid, and in guano islands the latter acid, if needed, can surely be imported to better advantage than prepared in this manner. The process, indeed, sins against that golden rule, "never use a nitrogenous substance as the raw material for a non-nitrogenous compound."

Prof. Wanklyn's process for the analysis of waters, which is introduced under the head, not of water, but of ammonia—is imperfectly described. The operator is directed first to distil the water *alone*, when the distillate will contain the free ammonia, and then to add the "alkaline permanganate and distil again for the determination of the nitrogen present in organic combination, or so-called albuminoid ammonia. A reference to Prof. Wanklyn's book on water analysis would have shown the writer that in order to find the nitrogen present in water in the form of ammoniacal salts it is necessary to distil the sample along with carbonate of soda. The writer afterwards mentions a solution of sodic carbonate, as "required for the application of the Nessler test," but he omits to say how and when such solution is to be used. The solution of chromic acid in water is described as being of a dark-brown colour. We have always found it of a scarlet.

Under the article "Beverages" we read: "Beer and porter are manufactured in enormous quantities in England, comparatively little being made anywhere else." Under "Alcohol," again, we find it stated that "the manufacture of ale and porter is confined to our own country." From these two passages the conclusion might be drawn that the production of malt-liquors was scarcely known abroad—an utterance at which Bavaria and Belgium would feel greatly astonished. At the foot of p. 193 we are told "it is reasonable to expect that a still further impetus will be given to the production of wines and spirits in England." On turning over the leaf we find that the author has also "turned over a new leaf," since he writes: "Wines . . . are not produced in any quantity in the British Isles."

In treating of hydrochloric acid, and generally of the noxious fumes arising from manufacturing operations, the authors quote some interesting statistical results obtained and published by the Belgian Government, showing that in four districts the total average death-rate had actually fallen since the introduction of chemical works. In one of the districts, however, there was an increase. We must beware, therefore, of laying too much weight upon these figures, especially as an apparently plausible attempt has been made to show that the health of a certain neighbourhood had been improved by the installation of a sewage-irrigation farm.

We shall watch the further progress of this work with much interest, trusting that as it proceeds its defects will be eliminated, and its many good features be rendered more prominent.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 6, August 11, 1879.

Experimental Researches on the Erosive Action of Gases, strongly Compressed and Heated, with Reference to the History of Meteorites.—M. Daubrée.—In this memoir the author shows that the erosive force of gases increases very rapidly with their pressure and temperature. To the purely mechanical phenomena there are always superadded calorific effects, and often chemical reactions. All these actions acquire a surprising energy when the gases, instead of whirling round in a space closed on all sides, are projected in a determined direction; for instance, when they escape by a narrow aperture.

Acids formed on the Distillation of the Crude Acids obtained by the Saponification of Neutral Fatty Bodies in a Current of Superheated Steam.—MM. A. Cahours and E. Demargay.—The acids formed under these circumstances are the valerianic, the caproic, cenanthylic, and caprylic.

Reply to the Remarks of M. Berthelot on the Author's Note concerning Chloral Hydrate.—A. Würtz.—The author defends his method of experimentation against the strictures of M. Berthelot.

Production of Electricity by the Rays.—C. Robin.—The author confirms by fresh experiments his former demonstration (A.D. 1865), that the electric organ in the tail of the Ray acts like those of the Torpedo and the Gymnotus, the differences being merely of a secondary nature.

Distillation of Liquids under the Influence of Static Electricity.—D. Gernez.—The distillation observed is almost exclusively a transportation of the liquid effected by means of electricity along the moistened (and therefore conductive) sides of the apparatus. The author finds, however, no relation between the quantities of the liquids thus conveyed, other things being equal, and the capillary constants of these liquids. There is a certain agreement between the manner in which the phenomena varies and the conductivity of the liquids. Doubtless if the liquid is a very bad conductor, like carbon, chloride, and sulphide, no distillation takes place, but if the charge passes at all the transference is the more abundant the lower is the conductive power of the liquid. If we submit to the action of the discharge a homogeneous mixture of two liquids, the action of the electricity determines a partial separation of the two liquids, which distil in proportions having no relation either with the volatility or the conductivity of the substance.

The Currents of Ampère, and on Residual Magnetism.—M. Tréve.—A shock upon an electro-magnet, whether solid or tubular, at the moment when the current ceases, is sufficient to diminish the duration of its demagnetisation to a considerable extent, and is consequently a remedy for remanent magnetism.

Vapour-Densities of Certain Organic Bodies Boiling at High Temperatures.—L. Troost.—The observed vapour-density of anhydrous phthalic acid is 5.23 (by calculation 5.13), and its volume equivalent is 4. Resorcin has the vapour-density 3.807 observed, 3.81 calculated. Ethyl-benzoic ether, 5.55 observed, and 5.2 calculated. Amyl-benzoic ether, 6.69 observed, and 6.65 calculated. The volume equivalent is in each case 4.

Density of Chlorine at High Temperatures.—A. Lieben.—The author comments on the fact observed by V. and C. Meyer that at temperatures exceeding 600° the

^{sp.} gr. of chlorine is only 1.63 instead of 2.45 (air = 1). He seeks to explain this fact by the supposition that above 700° chlorine follows a law of expansion different from other gases, its coefficient of expansion being possibly somewhat higher than those of oxygen, nitrogen, gaseous sulphur, &c. He suggests as another explanation that at high temperatures the molecules of chlorine, Cl_2 , are dissociated into isolated atoms.

Synthesis of Phenol-glucoside and of Ortho-formyl-glucoside or Helicin.—A. Michael.—The reaction of aceto-chlor-hydrate with potassium salicylate gives rise to helicin.

Compound of Chromic Acid with Potassium Fluoride.—L. Varenne.—The author assigns to this compound the formula KFl_2CrO_3 . It is resolved by alkalis into potassium fluoride and an alkaline chromate.

Production of Crystalline Magnetic Oxides by means of Potassium Cyanide.—L. Varenne.—A stannous salt treated with potassium cyanide precipitates stannous oxide, which, after being boiled with the cyanide for two or three days, is transformed into a crystalline oxide.

Identity of Hydrate of Di-isopren and Caoutchun with Terpen.—G. Bouchardat.—The nature of this paper appears sufficiently from its title.

No. 7, August 18, 1879.

Observations on the Reply of M. Würtz relative to Chloral Hydrate.—M. Berthelot.—Our learned colleague has not effected the justification, indispensable in every physical inquiry, of the limits of his experimental errors. The observation which he adduces relative to the rise of temperature produced in his apparatus by the union of nitrogen binoxide and oxygen cannot supply it, but may rather be invoked to show the whole extent of the causes of error inherent in his method.

Scintillation of the Flames of Gas Lamps.—F. A. Forel.—The author has observed that the flame of gas lamps when viewed from a considerable distance scintillates like the stars, and considers that the laws of this phenomenon may be thus conveniently studied, as all the conditions of the atmospheric stratum between the source of light and the observer, such as the pressure, moisture, temperature, &c., may be much more readily ascertained than similar conditions in the upper regions of the air.

Absorption of Nitric Oxide by Ferrous Salts.—J. Gay.—Ferrous salts absorb nitric oxide in variable proportions according to the temperature and the elastic force of the atmosphere of binoxide remaining in contact with the salt. Up to 8°, and under the atmospheric pressure, the quantity of nitric oxide absorbed by ferrous salts corresponds to 10NO_2 for 1 equiv. = 28 of iron. Hence, the formula of the compound is $3(\text{FeSO}_4) + \text{NO}_2$. Above that temperature and up to about 25° and under a pressure of about 760 m.m. the quantity of nitrous oxide absorbed is less, and corresponds to Peligot's formula, $4(\text{FeSO}_4) + \text{NO}_2$. Above 25° the formula of the saturated solutions agrees with $5(\text{FeSO}_4) + \text{NO}_2$. The author has succeeded in expelling all the nitric oxide and recovering the ferrous salt unchanged by passing through it a current of hydrogen, air being excluded. Reducing agents decompose the nitric oxide contained in ferrous salts, and liberate a mixture of nitrogen and nitrous oxide.

Reaction of Zinc Chloride upon Normal Butylic Alcohol.—MM. Le Bel and Greene.—The butylene of erythrite, our normal dimethyl-ethylen, is the final term of the reactions of zinc chloride upon the butylic alcohols.

Determination of Urea in Urines.—G. Esbach.—The author criticises the results furnished by M. Méhu on July 21. It is correct that a solution of glucose and of urea yields more gas than if the urea were alone, but the excess instead of being proportionate to the quantity of urea varies according to the glucose added.

Thermic Studies upon Nitro-glycerin.—H. Boutmy.—A thermic comparison between the direct process of manufacturing nitro-glycerin and that of Vonges, which turns on the mutual reaction of the sulpho-glyceric and sulpho-nitric acids.

Elimination of Bromine from Bromo-citraconic Acid and on a New Organic Acid.—E. Bourgoïn.—The author has eliminated the whole of the bromine from bromo-citraconic acid by means of silver oxide, a new acid being formed. This acid differs from the pyromucic by containing 2 equivs. of hydrogen more, and from citraconic by containing 2 equivs. of hydrogen less.

On Scandium.—P. Clève.

Oxygen Acids of Sulphur.—M. Maumené.—The action of iodine and barium hyposulphite, which yielded to Fordos and Gelis the tetrathionates, can yield according to the author's theory seven other acids. Two of these which precede S_4O_5 , the author has already obtained, namely S_2O_3 and S_6O_8 . The second of these is easily formed on mixing 3 equivs. of the hyposulphite with 2 of iodine.

Composition of Slate.—M. Maumené.—The author points out that certain kinds of this rock contain as much as 0.5 per cent of calcium carbonate, and are consequently readily attacked by atmospheric agents.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 63, March, 1879.

Report by M. le Comte du Moncel on M. Gaiffe's Peroxide of Manganese and Zinc Chloride Battery. The advantages of the Leclanché battery (manganic peroxide and sal-ammoniac) are well known, as it will act for entire years without attention. Hence it is almost universally applied for electric purposes where no great power is needed. If the manganic peroxide, however, is once exhausted they are rendered unserviceable, and must be replaced by new ones. M. Gaiffe, by his new arrangements, has overcome this difficulty, so that the elements may be charged as easily as in other batteries. He places the peroxide of manganese in several deep holes excavated in the cylinder of carbon, which forms the negative electrode and which serves at the same time as a porous vessel. This carbon is steeped in a solution of zinc chloride, which serves as the exciting liquid, and a rod of amalgamated zinc forms the negative pole. The chloride of zinc should contain from 15 to 20 of the salt of zinc, as neutral as possible and free from lead. The manganese should be in granules. The electromotive force of this element is equal to a couple and a half of Daniell.

Gazzetta Chimica Italiana.
Anno ix., 1879. Fasc. 1.

Various Observations on Digallic Acid.—Prof. Ugo Schiff.—A lengthy paper not susceptible of useful abstraction.

Reaction of Salicylic Acid with Salts of Iron.—Stefano Pagliani.—Robinet proposed a method for the detection of salicylic acid in wines and urine, which consists in treating the liquids with acetate of lead, filtering, adding to the filtrate an excess of sulphuric acid, and applying to the clear solution a few drops of ferric chloride, when a violet colouration shows the presence of salicylic acid. Marty and Dollfus call this process in question, affirming that the reaction is not produced in neutral ferric solutions, and is hindered by free acids, even the acetic. The author finds that the excess necessary to destroy the reaction by a known quantity of water containing sulphuric acid is when its weight is 400 times greater than that of the salicylic acid present. If nitric acid be used

in place of sulphuric the excess must not go beyond 385, and in case of hydrochloric acid the proportion must not be 36 times greater than that of the salicylic acid.

Action of Sulphuric Anhydride upon Pseudo-sulpho-cyanate of Phenyl.—G. Magatti.—The author obtained by this reaction a yellowish crystalline body of the composition $C_7H_5NS_2O_3$. On treatment with boiling water it is resolved into hydrosulphuric, carbonic, and sulphanic acids.

Preparation of Naphthyl Urea.—S. Pagliani.—The author has prepared and examined dinaphthyl urea.

Poisonous and Crystalline Ptomain extracted by means of Ether from two exhumed Bodies in which Arsenic had been Detected in Quantity.—F. Selmi.—The author describes the reaction of this compound, and which appear to be those of an alkaloid. It produced strongly-marked symptoms of poisoning in a frog, though the absence of arsenic and phosphorus had been previously ascertained.

Genesis of Poisonous Alkaloids in Corpses.—F. Selmi.—These compounds seem to be formed on the slow decomposition of albuminoids in the absence of oxygen.

Phenomena observed in the Plastering of Wines and Musts.—Prof. E. Pollacci.—The sulphate of potassa produced on plastering is an acid, not a neutral salt. If gypsum is added during fermentation the proportion of alkaline sulphate in the wine may reach 5 or 6 grms. per litre. Calcium tartrates and sulphates are further present in such quantity that the wines may be regarded as saturated solutions of these salts. In the fermentation of plastered grapes sulphuretted hydrogen is evolved, and ethylic mercaptan is formed, sometimes in quantity perceptible both by smell and taste.

Anno ix., 1879. Fasc. 2.

New Researches on Picrotoxin.—E. Paterno and A. Ogliaro.—The authors have discovered a new method for preparing picrotoxic hydrate in quantity. They heat an alcoholic solution of picrotoxin to ebullition, saturate with dry gaseous hydrochloric acid, distil off the bulk of the alcohol, mix the residue with water, and agitate the aqueous solution repeatedly with ether. The ethereal solution on evaporation leaves an abundant residue of picrotoxin hydrate. They have also obtained certain substitution products, such as a mono-benzoic and a bi-acetic.

Supposed Identity of Columbin and Limonin.—E. Paterno and A. Ogliaro.—These products differ in composition, in boiling-point, and in other characters.

Researches on Cinchonin.—M. Fileti.—An examination of the chlorine and bromine substitution derivatives of this base.

Analysis of the Spinell of Tiriolo in Calabria.—Dr. F. Mauro.—This mineral is a triple aluminate of zinc, manganese, and iron, along with a trace of antimonious acid.

Analysis of the Thermal Waters of Termini-Imerese.—E. Paterno and G. Mazzara.—Of no general interest.

Disulph-ethylidenic Acid.—Prof. J. Guareschi.—A description of this acid and of a number of its salts.

Gum of the Coloured Quebracho (*Lexispterigium Lorentii*).—Prof. N. Arata.—This exudation is of a red colour; it is soluble in boiling water, which, on cooling, deposits a part only; insoluble in benzol, carbon disulphide, chloroform, and essence of turpentine; but it dissolves readily in alcohol, acetone, and acetic ether. It is sparingly soluble in amylic alcohol and acetic acid. Hence it can neither be regarded as a gum nor as a true resin. It has no analogy with catechu or gambir, but approaches rather to kino.

Chemiker Zeitung.

No. 14, April 3, 1879.

Application of Watery Vapour in the Distillation of Liquids.—L. Ramdohr.—It is found that in the distillation of the liquid and solid hydrocarbons from lignite-tar, petroleum, rosin, rosin oils, &c., the introduction of superheated steam is very useful to prevent undesirable decompositions. The author has contrived an apparatus by which the steam is superheated inside the still, its temperature agreeing with the boiling-point of the contents.

Poisonous Colours.—According to the *Apotheker Zeitung*, of 118 samples of children's toys officially examined in 1878, 53, or nearly one half, were found adorned with poisonous colours. In 46 of these cases the vendors were punished.

Tannin in Algarobillo.—According to R. Godefroy (in the *Zeitschrift d. Oest. Apoth. Vereins*) the pods of *Balsamocarpum crevifolium*, or algarobillo, contain 68.38 per cent of tannin, and are an excellent material for the manufacture of this substance.

Die Chemische Industrie.

No. 7, July, 1879.

Injury to Vegetation by Acid Gases.—R. Hasen-clever.—The author admits that many causes in addition to the fumes of chemical works have had a deleterious action upon trees. He points out that the use of coal as fuel, whether in private houses, in mechanical manufactures, &c., has exerted a very devastating action upon vegetation. [The inference is, therefore, plain that restrictions upon the chemical arts to whatever length they might be carried must fail to prevent the injuries of which farmers and land-owners complain.] The memoir is accompanied by a coloured engraving of the leaves of various trees as affected by acid vapours. German observations confirm the view entertained in this country that the plane-tree is distinguished by its power of resisting acids.

Atti della R. Accademia dei Lincei.

Fascicolo 2, 1879.

Chemical Analysis of the Spinell of Tiriolo in Calabria.—Dr. F. Mauro.—Already noticed.

Disulphethylidenic Acid.—Prof. Guareschi.—Already noticed.

Secure and Delicate Process for the Toxicological Detection of Arsenic.—Prof. Selmi.—The method is that of Schneider modified so as to incur no losses. The substance to be examined is treated with hot concentrated sulphuric acid, and during the same time is traversed by a current of hydrochloric acid gas, which carries with it all the arsenic in the state of chloride, separating it from the organic substances with which it was mixed. The arsenical liquid is then placed in a Marsh's apparatus and tested in the usual manner. The author has been thus able to obtain the metallic ring on operating upon 100 grms. of animal matter containing 1-400th of a milligram of arsenious anhydride. The author criticises the process of Gautier, which answers for recent matter, but should not be adopted if the subject is putrid or mummified.

Fascicolo 3, 1879.

This part contains no chemical papers.

Reimann's Färber Zeitung.

No. 30, 1879.

Remarks on Pittical.—If pittical is separated from its alkaline salts, and heated with alcoholic ammonia in a sealed tube to 176°, a base is obtained closely resembling rosanilin in its composition. It is a triamin, and dissolves in acetic acid with a corn-flower blue colour. The new colour is the first member of a new series of tinctorial products, which may prove of great importance.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

AUGUST, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

	Appearance in a foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	An- Sulphuric hy- dride.	Hardness on Clark's Scale		
		Saline.	Organic.								Before Boiling.	After Boiling.	
													Grs.
<i>Thames Water Companies.</i>													
Grand Junction	Clear	0'000	0'011	0'120	0'120	20'20	7'750	0'518	1'008	1'367	15'2	5'0	
West Middlesex	Clear	0'000	0'011	0'120	0'149	20'90	8'060	0'468	1'008	1'267	15'7	4'8	
Southwark and Vauxhall	Clear	0'000	0'006	0'105	0'109	20'50	8'090	0'432	1'008	1'266	15'7	4'8	
Chelsea	Clear	0'000	0'010	0'090	0'101	20'00	8'010	0'396	1'080	1'330	14'7	5'4	
Lambeth	Clear	0'000	0'010	0'135	0'123	20'60	8'620	0'812	1'080	1'500	16'3	5'3	
<i>Other Companies.</i>													
Kent	Clear	0'000	0'002	0'420	0'028	29'00	8'170	0'696	1'800	1'400	19'8	5'8	
New River	Clear	0'000	0'004	0'120	0'107	20'20	8'170	0'460	1'008	1'430	15'4	5'0	
East London	Clear	0'000	0'008	0'111	0'073	20'70	7'630	0'648	1'008	1'400	15'2	5'8	

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

TO CORRESPONDENTS.

WE cordially thank those gentlemen who kindly supplied us with the requisite information relative to the various Examining Bodies, Colleges, and Schools of Chemistry given in our Students' number.

ERRATUM.—On page 125, line 4 from top, for Queenwood College, near Stockbridge, Hants, Mr. E. W. Prevost, Ph.D., read Queenwood College, near Stockbridge, Hants, Mr. H. Wilson Hake, Ph.D. The instruction at this college commences, we are informed, on the 25th inst.

J. Parry.—You will find an explanation in any text-book of Physics.

INSTITUTE OF CHEMISTRY.

The President has offered Two Prizes of £50 each for the two best original investigations involving Gas Analysis. These Prizes will be open to Associates, and to all persons (except Fellows of the Institute) who shall before the 31st December next have qualified for the Associateship in all respects short of passing the prescribed practical examination, and successful competition for these prizes will be accepted in lieu of such practical examination.—Further information may be obtained on application to the Secretary, Mr. C. E. GROVES, Somerset House Terrace, W.C.

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Midwifery and Diseases of ..	ALEXANDER SOMERS, M.R.C.S.
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peutics	CHAS. J. CULLINGWORTH,
Hygiene and Public Health ..	M.R.C.P. and S.
Medical Jurisprudence ..	JULIUS DRESCHFELD, M.D.,
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logy	DAVID LITTLE, M.D.
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Botany	

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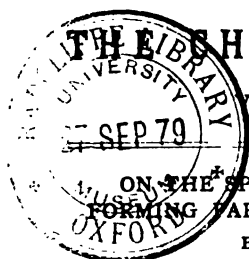
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CHEMICAL NEWS.

VOL. XL. No. 1035.

ON THE SPECTRUM OF THE EARTHS FORMING PART OF THE YTTRIA GROUP.

By M. J. L. SORET.

M. Clève communicated to the Academy at their Meeting on Sept. 1, a note on two new elements in erbia (see CHEMICAL NEWS, vol. xl., p. 125). I crave permission to offer the following remarks on that note.

(1.) M. Clève characterises one of those two elements, for which he proposes the name of holmium, by two absorption rays, one in the red, $\lambda=640$, the other in the yellow-green, $\lambda=536$. It doubtless escaped his memory that more than a year ago* I pointed out those two rays as not belonging to erbia, but to a new earth, whose probable existence had been announced by MM. Delafontaine and Marignac, and which, for the time being, I designated X. Since then M. Delafontaine has given the name of *philippia* to a substance he has pointed out as identical with X. The description which he has at present given† is not sufficiently complete to enable me to deny or affirm this identity, but I am inclined to admit it, taking into account, however, some reservations made by M. Delafontaine on the purity of his product.

I would remind you also that besides the two above-mentioned rays, I recognised in the visible spectrum of the earth X (solar light) three other rays or absorption-bands, the one less refrangible than A, the second overlapping the ray of erbia, $\lambda=450$, in the indigo;‡ the third (faint) in the violet, a little beyond h. Finally, I described the ultra-violet spectrum of this earth, which is still more characteristic, and shows six maxima of absorption from H to R.

Since that time these results have been confirmed by the examination of a great number of products placed at my disposal by M. Marignac. I have constantly found that these different rays increase or diminish in intensity simultaneously, whilst the erbia rays behave quite differently.

In samarskite this earth X is, relatively to erbia, much more abundant than in gadolinite. Thus, in the majority of the products of samarskite, the ray $\lambda=536$ is darker than the ray α of erbia, $\lambda=523$. It is habitually the reverse in the gadolinite products.

I do not think that the difference between the rays of erbia and those of the earth X can be explained by actions analogous to those observed by Prof. Lawrence Smith and M. Lecoq de Boisbaudran in their researches on the spectra of didymium and erbium nitrates, when an excess of acid was added.§ I have, however, found the same general characteristics in the chlorides and nitrates with excess of acid.

Upon the whole I think the existence of the earth X is fully demonstrated, but I see in M. Clève's note no result proving holmium to be a different body.

(2.) M. Clève characterises by a red ray $\lambda=684$, his second new element, which he proposes to call *Thulium*. I have already directed attention to the fact that in the

products rich in earth X, and at a feeble equivalent this ray 684 is not to be met with, whilst all the other rays of erbia can easily be distinguished.* I added in a note that the variations of this ray were perhaps in correlation with some new facts observed by M. Marignac, who, indeed, was then occupied in the preparation of ytterbia, and had given me some products to examine with the spectroscopic, in which products the latter earth was already strongly concentrated. In these products the ray 684 had taken a predominant intensity, whilst those of erbia, and especially those of the earth X, were weakened. But when the ytterbia had been still further purified, the intensity of the ray 684 diminished also, and in the last product it existed only as a trace, as M. Marignac has stated in his paper.† The following results were, for example, obtained on the four principal rays of the red, which formerly were attributed to erbia.

Rays visible in the Order of their Intensity.

Ytterbia, almost pure ..	Equivalent 131: only a trace of ray 168 is to be seen.
Mixture	Equivalent 128: 684 (well-marked; 653 (faint); 667 (640 no trace).
Mixture	Equivalent 124: 653 (well marked), 684, 667, 640.
Mixture	Equivalent 120: 653, 684; then 667 and 640 perceptibly equal.

If, then, neither I nor M. Marignac, who would have been far more authorised in doing so, have drawn no positive conclusion from these facts, it is because it seemed to us that in a question so difficult it is perhaps premature to affirm the existence of a new element whilst it is still impossible to isolate it or to determine its chemical characteristics, the only ground for the supposition being the presence of a single ray in the absorption spectrum.—*Comptes Rendus*, September 15, 1879.

RESEARCHES ON ERBIA.

By M. LECOQ DE BOISBAUDRAN.

In the *Comptes Rendus* for September 1st M. Clève announces that he has divided erbia into several distinct earths, which he proposes to name *Erbia*, *Oxide of Thulium*, and *Oxide of Holmium*. The absorption spectrum attributed originally to erbia alone would thus be the result of the superposition of the spectra of three earths.

It is to be remarked that the holmium rays are precisely those pointed out by M. Soret as most characteristic of his earth, X. The two substances are evidently identical.

I am ignorant of the extent to which M. Soret has carried his examination of the red ray attributed by M. Clève to thulium, but he had specially pointed out to me its existence (to the almost entire exclusion of the other red rays of erbia) in a sample of impure ytterbia which he sent me a few months ago. This ray always seemed to me much less narrow than that of ordinary erbia, and appeared to form rather a broad band.

After a visit paid by M. Soret to M. Wurtz's laboratory, where I was working last spring, I undertook some experiments on the question of the plurality of erbia earths. Detained as I am at Cognac, and consequently prevented from immediately completing the examination of my products, left behind in Paris, I must beg the Academy to receive the following statement of my observations, incomplete as they still are.

On M. Soret's announcement of the probable existence

* See *Comptes Rendus*, April 29, 1878 (CHEMICAL NEWS, vol. xxxviii., p. 237). The paper is published in *extenso* in the *Archives des Sciences Physiques et Naturelles*, vol. lxiii., pp. 99 and 101.

† *Comptes Rendus*, October 14, 1878; CHEMICAL NEWS, vol. xxxviii., p. 202.

‡ It is this band which M. Delafontaine points out as characteristic of *Philippium*. I consider that indication an unfortunate one, for, according to all the observations I am acquainted with, erbium gives rise to a much narrower ray at the same place.

§ *Comptes Rendus*, June 9, 1879; CHEMICAL NEWS, vol. xxxxi., pp. 286, 298.

* *Archives des Sc. Phys. et Nat.*; loc. cit., p. 99.

† *Archives des Sc. Phys. et Nat.*, 1878, t. lxiv., p. 101. M. Marignac also pointed out some other reasons which would lead us to suppose that these products were complex mixtures.

of at least two distinct oxides in erbia, I had examined the absorption spectra of salts of erbium of different composition. All these compounds, including those extracted from euxenite (a mineral chemically analogous to samarskite), but not those obtained from samarskite, gave the same rays and the same relative intensities as the chloride of erbium used for my old diagram (*Spectres lumineux*, Pl. xv.). The salts of erbia obtained from samarskite by M. Demarçay and Prof. Lawrence Smith present indeed the rays of my drawing, but with a marked alteration of relative intensities. The chief characteristic of samarskite-erbia is that—First, the green ray, $\lambda=536.3$, is much more intense than its neighbour, $\lambda=540.9$, whilst in the other erbias the predominance of $\lambda=536.3$ is very slight; second, the red ray, $\lambda=640.4$, is as strong, or even stronger, than its neighbour, $\lambda=653.4$, whilst in the other erbias $\lambda=653.4$ predominates greatly over $\lambda=640.4$. The other variations of intensity are less striking.

If ordinary erbia is a mixture of several analogous earths, is it not strange to see specimens of very different productions possessing a composition, the constancy of which is shown by the identity of the spectra? On the other hand, the presence of foreign bodies, the nature and even the quantity of the acids have often a very marked influence on the absorption-bands. We might then, strictly, still admit the unity of erbia, and suppose that the special distribution of intensities in the spectrum of samarskite-erbia is due to the presence, in samarskite, of a larger proportion of some principle, known or unknown, different from the earths properly so-called, but still undetermined.

These objections having been submitted to M. Soret, that eminent physicist gave reasons in favour of the existence of his earth X, which led me to undertake the following research:—

I selected two rich samples of erbia. One, A, almost pure, prepared by M. Demarçay, gave with great exactitude the spectrum of my drawing. The other, B, was a raw product, rich in yttria, obtained by Prof. Lawrence Smith's labours on samarskite, of which it presented the characteristic spectral type.

A and B were separately submitted to fractionated precipitations with ammonia. One of the extreme products of A soon showed a tendency to resemble the spectral type of B, whilst one of the extremes of B in like manner approached the type of A, the direction of the modifications being the same in the two series.

With ammonia the operation worked very slowly, but by the methodical use of the potassium and sodium sulphates in the cold, and by the aid of heat, the effect was analogous, and far more rapid. A few operations were sufficient to extract from A an erbia spectrally similar to that of samarskite, and from B an earth possessing a spectrum almost exactly like that of my old drawing.

These results seem to decide the question in favour of M. Soret's ideas, and agree with the important researches recently published by MM. Clève and Thalén. Moreover, before my opinion on such a delicate question is settled, I should like to wait for fuller information, and the completion of the work begun, confining myself at present to publishing the above simple facts of observation.—*Comptes Rendus*, September 15, 1879.

ON THE CHEMICAL COMPOSITION OF A NODULE OF OZOKERITE FOUND AT KINGHORNNESS.*

By W. IVISON MACADAM, F.C.S., &c.,
Lecturer on Chemistry, Edinburgh.

THE nodule to which this paper refers was discovered at Kinghornness during the excavations rendered necessary

by the fortifications at present being raised for the defence of the Firth of Forth. The material was enclosed in a nest or nodule, and was found at a depth of 15 feet from the surface of the ground, and embedded 5 feet in hard trap rock. The rock in which the nodule was obtained was sound, no crack or fissure being observable for several feet round the nest. At a point some distance below the nodule the section shows a series of small veins or fissures running through the rock in various directions, and averaging $\frac{1}{2}$ of an inch in breadth. The analysis of this rock gave the following results, calculated to percentages.

(1.) On treating the pulverised sample with hydric chloride (HCl), and subjecting the mixture to a prolonged low heat, it was found that 29.73 per cent of the substance dissolved in the acid. The detailed results of the analysis of this solution are as follows:—

	Per cent.
Ferric oxide (Fe_2O_3)	11.45
Aluminic oxide (Al_2O_3)	3.63
Calcic oxide (CaO)	3.70
Magnesian oxide (MgO)	0.37
Potassic oxide (K_2O)	0.13
Sodic oxide (Na_2O)	
Carbonic anhydride (CO_2)	8.17
Sulphuric anhydride (SO_3)	0.21
Silica from soluble silicates (SiO_2)	2.07
Total soluble in acids	29.73
Insoluble in acids and silicates	70.27
	100.00

(2.) The portion insoluble in acids was then fused with a flux, and the following results obtained from the after solution in acids:—

	Per cent.
Ferric oxide (Fe_2O_3)	9.92
Aluminic oxide (Al_2O_3)	5.36
Calcic oxide (CaO)	8.17
Magnesian oxide (MgO)	5.67
Silica from silicates, &c. (SiO_2)	41.12
Total from fused portion	70.24
Soluble in acids	29.73
	99.97

The rocks lying next to the trap were also analysed, and gave results as stated below:—

1. Soluble in Hydric Chloride (HCl).

	No. 2. Rock above Trap.	No. 3. Rock below Trap.	No. 4. Same as 2, but nearer the surface ground.
Ferric oxide (Fe_2O_3)	21.37	31.48	15.78
Aluminic oxide (Al_2O_3)	4.06	4.59	7.92
Calcic oxide (CaO)	2.83	3.11	2.19
Magnesian oxide (MgO)	2.98	4.23	2.47
Potassic oxide (K_2O)	0.98	0.82	1.48
Sodic oxide (Na_2O)			
Carbonic anhydride (CO_2)	5.19	6.01	4.62
Sulphuric anhydride (SO_3)	0.32	0.28	1.62
Sulphur (S)	0.13	0.16	2.17
Silica (SiO_2) from sol. silicates	2.08	2.23	3.95
Moisture	3.22	3.41	3.14
Total soluble in acid	43.16	56.32	44.34
Insoluble in acid	56.84	43.68	55.66
	100.00	100.00	100.00

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

II. Matter Insoluble in Hydric Chloride (HCl) fused with Flux and then treated with Acid.

	No. 2. Rock above Trap.	No. 3. Rock below Trap.	No. 4. Same as 2, but nearer the surface of ground.
Ferric oxide (Fe_2O_3)	5.28	4.84	7.04
Aluminic oxide (Al_2O_3)	3.48	4.12	7.76
Calcic oxide (CaO)	2.15	1.18	0.78
Magnesian oxide (MgO)	1.03	0.28	0.24
Silica (SiO_2) from silicates, &c.	44.68	33.12	39.72
Total from fused portion	56.62	43.54	55.54
Soluble in acids	43.16	56.32	44.34
	99.78	99.86	99.88

The nodule when broken consisted of—

- (1.) An outer coating of hard rock;
- (2.) An inner lining of calcite crystals; and
- (3.) Centre nodule of bituminous matter.

When first brought to light the calcite crystals were almost black in colour, due to a certain amount of the bituminous matter, but this slowly evaporated, and left the crystals pure white in colour. The analyses of these crystals of calcite yielded the following results:—

	Black Crystals containing Bitumen.	White Crystals.
Calcic carbonate (CaCO_3)	96.76	98.11
Ferrous carbonate (FeCO_3)	0.19	0.21
Magnesian carbonate (MgCO_3)	0.31	0.33
Silica (SiO_2)	1.06	1.22
Bituminous matter	1.68	0.13
	100.00	100.00

The lower veins or fissures were also calcite-lined, and contained within this coating the bituminous matter.

Besides the nodule found at Kinghornness, another nodule of a similar character was obtained on the Island of Fuchkeith, embedded in solid trap, 10 feet from the surface, and a small spring of water on that island smells and tastes distinctly of paraffin products.

The Kinghornness nodule has a distinct bituminous odour; is a lustrous black; amorphous, soft solid, easily cut with the nail, and pliable between the fingers. The specific gravity is 970 (water 1000), so that the nodule floats upon water. It fuses at 176° F., and becomes solid on cooling. Experiments with the various solvents upon the bituminous material showed that water and ordinary acids had practically no action whatever. Alcohol, both hot and cold, had a very slight solvent power, but ether dissolved a considerable proportion, giving a brown solution, and turpentine readily acted upon the substance, yielding a deep brown-black solution. The ethereal liquid had a fine iridescent green colour when viewed by reflected light. The substance of the nodule readily burns when lighted, giving a strongly luminous flame. The analysis of the contents of the nodule gave as follows:—

	Per cent.
*Volatile organic matter	99.38
Ash or mineral matter	0.61
	99.99

Volatile gaseous matter given off below 212° F.=5.961 per cent.

When distilled at a bright cherry-red heat, 26.57 grs. being used, the results gave—

	Grains.
Volatile matter	15.93
Coke, { Fixed carbon	10.48
10.64 grs. { Ash (mineral matter)	0.16
	26.57

These results calculated to the percentages give—

	Per cent.
Volatile matter	59.955
Coke, { Fixed carbon	39.431
40.041 p.c. { Ash (mineral matter)	0.610
	99.996

The coke left behind after this treatment was a hard, black, shining, porous mass, and the ash obtained by incinerating it was pure white, and principally consisted of calcic carbonate (CaCO_3) and silica (SiO_2).

The material was afterwards submitted to destructive distillation at a black-red heat, when the substance was found to split up into six distinct parts—four distillates, a coke, and a volatile non-condensable gas. In this operation 58.29 grains were used, and yielded—

	Grains.
First distillate	5.65
Second distillate	11.35
Third distillate	21.33
Fourth distillate	14.72
Total volatile condensable products	53.05
Coke	2.28
Uncondensable gases	2.96
Total	58.29

Calculated to percentages the results stand—

	Per cent.
First distillate	9.694
Second distillate	19.471
Third distillate	36.592
Fourth distillate	25.253
Volatile condensable products	91.010
Coke	3.911
Uncondensable gas	5.078
	99.999

When first heated the substance fused and frothed, and on the further application of heat gave the first distillate, which was of a grey colour, somewhat mobile, and had a disagreeable odour. The second distillate was black in colour, fully more mobile than the first distillate, and also possessed a most disagreeable odour. The third portion was more mobile than the second, had a yellow colour and a marked paraffin odour. The fourth distillate was obtained after raising the heat, had a yellow-red colour, was liquid whilst hot, but turned solid on cooling, and had also a paraffin odour. The uncondensable gaseous matter readily burned on the application of a white light, gave a pale non-luminous flame, and possessed all the chemical properties of methane or marsh-gas (CH_4). The carbon left in the retort, added to the amount of uncondensable gas, gives 8.989 per cent, and on calculating the uncondensable gas into ethylene or olefiant gas (C_2H_4) the result obtained is 8.886, or nearly identical.

These results go far to show that the bituminous-like matter of the nodule consists of a member or members of the olefine (C_2H_4) series of organic compounds, a point which is further strengthened by the fact that the carbon and hydrogen in the original substance are contained therein in almost exactly the necessary proportions to form an olefine.

It is probable that the source of the contents of the nodule lies in one of the coal or shale beds abounding in the district, and that a low internal heat has dissolved the material from its parent stratum. That the heat was low, or certainly not above a cherry-red, is certain, else the olefine would have been split up into a member of the methane or CH_4 group of organic substances, accompanied by a deposition of free carbon.

ON THE
SEPARATION OF PHOSPHORUS AND IRON,
ESPECIALLY WITH REFERENCE
TO THE MANUFACTURE OF STEEL.*

By THOMAS BLAIR.

ALL recent improvements in the manufacture of iron and steel sink into comparative insignificance before the successful production of merchantable steel from phosphoretic pig-iron, and it has been thought advisable that the writer should prepare a short review of the whole subject of the purification of iron from phosphorus.

Pig-iron being in nearly all cases the raw material from which iron or steel in the finished state is prepared, it is necessary to know the amount of phosphorus it contains, its suitability for various purposes being dependent upon this; and, as a necessary consequence, the selection of ores for smelting depends mainly on the same fact, viz., their content of phosphorus.

In the following table the percentage of phosphorus existing in ores of various kinds is given, the first two being suitable for the manufacture of steel and fine wrought-iron, the others not:—

	Iron Per cent.	Phosphorus Per cent.
1. Spanish hæmatite	50·63	0·01
2. Cumberland hæmatite.. ..	58·00	0·01
3. Cleveland ore (raw)	30·00	0·50
4. Northamptonshire ore ..	39·00	0·40
5. Lincolnshire ore	30·00	0·35 to 0·40
6. Clay-band, or argillaceous ironstone	30·00	0·20

In the pig-iron made from such ores, the amount of phosphorus may be previously estimated by calculating the quantity of ore required to produce a ton of pig-iron. The following gives the percentages of phosphorus usually found in pig-iron smelted from the above ores:—

1. Spanish pig	0·03 p.c.
2. Cumberland pig.. ..	0·03 "
3. Cleveland pig	1·50 "
4. Northamptonshire pig ..	1·30 "
5. Lincolnshire pig	1·25 "
6. Clay-band, or "Mine" pig ..	0·50 "

In the products of purification, the following are the percentages of phosphorus usually found:—

	Phosphorus per cent.
1. Refined iron	0·35
2. Puddled iron (common).. ..	0·08
3. " " (best)	traces
4. Crucible steel	trace to 0·03
5. Bessemer or Siemens's (rails) ..	0·05 to 0·20
6. " " (soft)	0·02 to 0·06

It will be obvious from these figures that, if the purifying processes are unable to deal rapidly and energetically with phosphorus, only such pig-irons as Nos. 1 and 2 in the above table can safely be employed, since the presence of phosphorus in steel or wrought-iron, in quantities greater than the above, unfit them more than does any other element for most of the purposes for which they are intended, inasmuch that its presence in excess renders the metal hard and brittle, and although its malleability does not appear to be affected, yet in the cold it is exceedingly brittle. These faults are intensified when the other elements usually existing in wrought-iron or steel are present in larger proportions than usual. Mr. Hackney states that "Steel for rails if nearly free from phosphorus, and at the same time containing little silicon, is not made too hard to be safe by even 0·6 or 0·7 per cent of carbon; yet if the phosphorus amounts to 0·05 or 0·06 per

cent, the carbon cannot be prudently allowed to exceed 0·45 per cent, and steel rails containing 0·1 per cent of phosphorus should never contain more than 0·3 per cent of carbon, and so on until when the carbon is kept down to 0·16 per cent. Steel containing 0·3 per cent of phosphorus will yet make serviceable rails, and that containing 0·15 per cent will make excellent tough boiler-plates, that may be bent double cold after heating to redness and quenching in water; while, of course, metal free, or nearly free, from phosphorus is still softer and tougher, more, in fact, like copper than any commonly known variety of iron."

It will thus be seen that only those pig-irons in which the phosphorus is considerably less than 0·1 per cent can be employed in the manufacture of steel in the Bessemer, Siemens, or crucible processes, in which no elimination of this objectionable element takes place.

Unfortunately the bulk of the iron ores raised in this country, as well as in most other iron-producing countries, or in districts adjacent to coal-fields, contain so much phosphorus as to unfit them for the production of iron suitable for these just-named processes, such crude iron being smelted from purer and scarcer ores, which are often more expensive to raise, or which occur at great distances from iron-producing centres.

It would naturally occur to many that it might be feasible to separate the phosphorus from the ore previous to smelting; this has actually been done. Such processes are based upon the following chemical reactions:—Phosphorus exists in iron ores, either as lime or iron phosphate. Lime phosphate is soluble in a strong solution of sulphurous acid, but is not acted upon by ammonium sulphide. Iron phosphate is decomposed by ammonium sulphide, but is not soluble acid, and is also rendered soluble in water when fused with common salt.

Some years ago M. Jacobi, manager of Kladno Works, Prague, Austria, succeeded in extracting the lime phosphate very completely from ores in which it existed in quantity, by treating the coarsely-powdered ore with aqueous sulphurous acid; on boiling the solution the acid was recovered and re-condensed in coke towers, whilst the phosphate of lime was precipitated. This substance was said to fetch a price which amply repaid the cost of the process. But when it is remembered that a modern blast-furnace consumes from 600 to 1000 tons of ore per week, it will be evident that the magnitude of the plant required will be an effectual bar to its adoption; and that, further, as the ore treated in such processes requires to be pulverised, only a small proportion of the ore so treated could be used in the blast furnace.

In the earlier days of iron smelting, the little Catalan forges, still used here and there on the sides of the Pyrenees, had but imperfect powers of reduction, and large quantities of oxide of iron passed off into the slag, carrying away the whole of the phosphorus, so that iron of excellent quality was directly produced from very impure ores. The same ores smelted in a blast-furnace, where the powers of reduction are perfect, gave irons rich in phosphorus, with no waste of iron in the slags. In fact the blast-furnace is so perfect in its reducing powers that it reduces all the undesirable elements, which it is the object of subsequent oxidising processes to eliminate, the smelting process being essentially one of reduction, whilst all purifying processes are necessarily oxidising.

Dr. Percy aptly describes the blast-furnace process as "the indirect extraction of iron in the state of cast-iron, from the ore." Mr. I. Lowthian Bell thus tersely describes the theory of smelting:—

"Exposed to the intensely deoxidising agency of the blast-furnace, portions of the silica, and probably the greater portions of the sulphur compounds, lose their oxygen, and are taken up wholly or in part by the reduced iron. Practically, however, it may be assumed that all

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

* This statement, however, seems open to question, since extra soft steel seldom contains more than 0·06 per cent of phosphorus.

the phosphoric acid in the minerals is decomposed during the smelting process, and in consequence all its phosphorus is to be found in the pig metal. In the metallic product of the blast-furnace, carbon, derived either directly from the fuel, or from dissociated carbonic oxide, is invariably present. For our present purpose we may therefore regard pig-iron as a compound of carbon, silicon, sulphur, and phosphorus, and the metal itself."

Experiments made by Mr. Stead seem to prove that phosphorus exists in pig-iron as phosphide (Fe_3P) in a state of solution, or intimate diffusion throughout the mass, he having actually squeezed out a compound very rich in phosphorus from viscous pig-iron, an examination of which proved this to be the state of combination.

The prospects of successful removal of phosphorus in the blast-furnace are very discouraging. Attempts have been made to flux it off by adding salt, magnesian limestone, and other bodies, but without the smallest measure of success. Low temperatures and presence of oxide of iron in the slag being essential conditions for its removal, it is obvious that the very perfection of the blast-furnace renders phosphorus elimination impossible. When these conditions do occur, that is in disordered working, some phosphorus is actually slagged off.

Unpurified pig-iron is used only for castings, of which however, an enormous tonnage is annually made. It is stated that for small articles, where no great amount of strength is required, phosphoretic pig is more suitable, owing to the greater fluidity which phosphorus confers upon it. For all other purposes, pig-iron has to undergo some operation of refining in order to arrive at a degree of strength, which could only be attained with cast-iron by employing cumbersome masses.

Up to the time of Bessemer's invention, the puddling process was the principal one in use for this purpose. Before the invention of this process by Henry Cort, exactly a century ago, malleable iron was only obtainable in small quantities by wasteful and expensive processes, similar to the Catalan, Corsican, or Osmund. All of them depending for their success upon the fact that when molten iron is kept sufficiently long, and at a proper temperature, in intimate contact with oxide of iron, the whole of the phosphorus is removed, after the silicon has gone, and before, concurrently with, or after the carbon, according to circumstances to be hereafter explained.

It was formerly the practice to treat all pig-iron intended for puddling in a refinery, or "running-out" fire, as a preliminary to puddling. This is now, however, to a very great extent discontinued, except for the manufacture of special qualities of wrought-iron. Its chief merit appears to be that it removes most of the silicon. The process consists in melting under coke, by means of a blast of air impinging at an angle of about 45° , about one ton of pig-iron; oxide of iron being added in the shape of hammer or roll-scale. About one-and-a-half hours is the time occupied in melting; another half-hour is occupied in blowing air upon the bath of molten metal.

[A diagram accompanies the author's paper, in which was given the curves representing the removal of carbon, silicon, and phosphorus from the molten metal; the abscissæ representing times, and the ordinates percentages of the elements removed. This diagram is based on the following analyses given by Mr. I. Lowthian Bell in his paper on this subject, read before the Iron and Steel Institute in 1877:—

	Carbon Per cent.	Silicon Per cent.	Phosphorus Per cent.
Bowling cold-blast pig..	3.686	1.255	0.565
After melting in the refinery	3.510	0.575	0.557
10 minutes after fusion	3.707	0.478	0.537
20 "	3.644	0.273	0.530
28 "	3.544	0.154	0.509
Refined metal..	3.342	0.130	0.490]

The refined plate, or as is now most commonly the case, the crude pig-iron, is charged into the puddling furnace in

quantities of about 4 cwt. (the modern mechanical furnaces take quantities up to one ton), the bottom and sides of the furnace being previously lined with some variety of oxide of iron. In about half an hour the charge is melted, and the molten metal soon commences to boil, owing to the reaction between the oxide of iron and the carbon, jets of carbonic oxide gas being projected through the bath, burning with a blue flame on the surface, and a considerable quantity of slag is formed, consisting of ferrous ortho-silicate. The bath is diligently stirred by the puddler, or in the case of mechanical furnaces by the rotation of the furnace, in order to effect intimate contact between the molten iron and the oxides. As the carbon disappears, the iron becomes less and less fusible, and is subsequently removed in several balls of 80 to 100 pounds each, which are then forcibly compressed by hammering or squeezing to expel the intermingled slag and to consolidate the mass.

It will be well here to examine with care what takes place in these two processes, in order to understand the chemistry of the elimination of phosphorus in the Bessemer process.

First, the oxygen of the oxides, or of the air, or both, rapidly removes the most readily oxidisable element, silicon. Phosphorus is oxidised and passes off into the slag, most probably as tribasic phosphate of iron. Percy suggests that its removal takes place by liquation, or "sweating out," when by removal of the carbon the iron begins to thicken, or "comes to nature;" but the experiments of Bell, Stead, and others point conclusively to the fact that molten oxide actually washes out the phosphorus, oxidising it at the same time. It is worthy of notice that if the slag is not tapped out of the furnace the phosphorus is reduced and passes again to a considerable extent into the iron at the end of the operation.

The following are analyses of slags from the refinery and puddling processes:—

	Refinery p.c.	Puddling p.c.
Iron	50.96	49.65
Silica.. .. .	25.77	29.60
Phosphorus	1.37	3.50

It may be here noted that the addition of fluor-spar with titaniferous ore (known as Henderson's process) seems greatly to expedite the removal of phosphorus during puddling.

Bessemer Process.—In 1856 Sir Henry (then Mr.) Bessemer read a paper before this Association at Cheltenham, on a new process for manufacturing steel cheaply on the large scale. Up to this time the bulk of the steel made in the world was produced by the crucible method, by melting cemented, or carbonised puddled Swedish iron with the addition of manganese. His process has made such rapid progress that now the great majority of the steel made in the world is produced by its means, and it is generally so well understood that it requires a very brief description here.

Pig-iron of the following approximate composition:—

Graphite	2.90 p.c.
Combined carbon	0.50 "
Silicon	2.00 "
Sulphur	0.03 "
Phosphorus	0.05 "
Manganese	0.60 "

melted in cupolas or in reverberatory furnaces, or run direct from the blast-furnace, is run into a movable converter made of boiler-plate iron, lined 9 inches or more with gannister, and mounted on trunnions, so as to move through about three-quarters of a revolution by hydraulic machinery. When in position for blowing, the tuyeres are at the bottom of the vessel, and numerous jets of air are forced through the bath of metal at a pressure of from 15 to 20 lbs. to the square inch, according to the depth or liquidity of the bath.

As in the puddling process, the silicon is rapidly re-

moved, and subsequently the carbon. The phosphorus in the steel, however, is slightly more than that in the pig-iron used, owing to the fact that there is some waste of iron and none of phosphorus. A more concentrated solution may be said to be formed.

In a period of from ten to about thirty minutes, according to quantity treated, and proportion of silicon, carbon, or manganese present, the carbon is reduced to 0.1 per cent or less. The flame then "drops," or suddenly diminishes in size, while at the same moment the lines in the spectrum, due to the oxidation of carbon, are seen to suddenly disappear. The product is then, if cooled, a soft metal, of a composition very closely approximating to that of wrought-iron, but so spongy that it is valueless. By the addition of an alloy of iron, carbon, and manganese, well known as spiegeleisen or ferro-manganese, the dissolved or occluded gases are taken up, and a sufficient quantity of carbon and manganese restored to the bath to render the resulting steel hard or soft as desired. It is only right to add here that this necessary addendum to Bessemer's brilliant process originated with R. F. Mushet.

Owing to the rapid oxidation of silicon, carbon, and manganese the temperature and fluidity of the bath rapidly increase. The initial temperature of the molten pig being 1500° C., that of the steel, as poured, is about 2500° C., and a piece of thick platinum wire rapidly fuses in a stream of the steel.

It is now a matter of history that Sir Henry Bessemer avowedly thought at first that any pig-iron was suitable for his process, and that he had ultimately to fall back upon pig almost free from phosphorus.

In the Siemens, and Siemens-Martin process also, which there is not time now to describe, such non-phosphoretic irons are also a necessity. Thus both these eminently useful processes preclude the use of those irons made from the bulk of native ores, which, as already stated, contain phosphorus in notable quantity. The elimination of this element in these two processes, which produce the bulk of steel made in the world, was therefore always a necessity, but became a burning question when the iron rail trade died, bequeathing to the iron masters mountains of ore, hosts of idle blast- and puddling-furnaces and rolling-mills, the owners of which had no alternative but to contend with and overcome, if possible, that perverse but ubiquitous principle of nature which renders the attainments of objects most desirable extremely difficult.

The first to step into this ugly breach was Mr. Isaac Lowthian Bell, who gave to the world in 1877 the outcome of his careful study, and experiments on the separation of carbon, silicon, and phosphorus, as effected in the processes already described.

Mr. Bell's efforts were directed towards the attainment of a practical method of separating the phosphorus from Cleveland iron to such an extent as to permit of its use in the manufacture of steel, and also to make puddled iron of a quality superior to that obtained by the ordinary puddling process.

Pig-iron taken direct from the blast-furnace was treated in a suitable apparatus with melted oxide of iron, and it was found that whilst 11 per cent only of the carbon was removed, as much as 91 per cent of the phosphorus went; or in exact figures the pig-iron contained—

Carbon	3.637 p. c.
Phosphorus	1.351 "

The average composition of four samples so purified was—

Carbon	3.227 p. c.
Phosphorus	0.109 "

By prolonging the operation the whole of the carbon can be also eliminated, but that there is a slight increase in the phosphorus at the end.

This purified metal, after puddling, piling, and rolling, gave a finished bar containing—

Silicon	0.079 p. c.
Sulphur	0.007 "
Phosphorus	0.040 "

This analysis corresponds to that of a finished bar of high quality.

It may be here noted that Mr. Stead had on the small scale found that Cleveland iron was purified from its phosphorus, to a great extent, by violently shaking it with fused oxide of iron in a closed crucible. Mr. Stead also over-blew pig-iron in the Bessemer converter, but owing to the fact that his lining was silicious instead of basic, his experiment was unsuccessful.

One of Mr. Bell's proposals was to use the Bessemer converter as a means of rapidly removing silicon and then puddling the product; another was to make use of the Godfrey-Howson puddling-furnace in which to treat molten pig-iron with fused oxide of iron, or fused iron slags or iron ores.

The phosphorus removed from the iron naturally passes into the slag formed, so that oxide or slag containing 1.5 per cent of phosphorus before the operation, was found to contain 3 or 4 per cent afterwards.

The most noteworthy disadvantage of Mr. Bell's process, as compared with that of Thomas and Gilchrist, to be hereafter described, seems to be that it would only be capable of dealing with small quantities at a time, and that a very extensive plant would be required; whilst the other deals with 6 or 8 tons at one operation, and only requires slight modifications of the ordinary Bessemer plant.

(To be continued.)

THE ACTION OF HEAT IN VACUO ON METALS.*

By T. A. EDISON.

IN the course of my experiments on electric lighting I have developed some striking phenomena arising from the heating of metals by flames and by the electric current, especially wires of platinum, and platinum alloyed with iridium. These experiments are in progress.

The first fact observed was that platinum lost weight when heated in a flame of hydrogen, that the metal coloured the flame green, and that these two results continued until the whole of the platinum in contact with the flame had disappeared. A platinum wire four-thousandths of an inch in diameter, and weighing 306 m.grms., was bunched together and suspended in a hydrogen flame. It lost weight at the rate of a fraction less than 1 m.grm. per hour as long as it was suspended in the flame. When a platinum wire is stretched between two clamping posts, and arranged to pass through a hydrogen flame, it is coloured a light green; but when the temperature of the wire is raised above that of the flame, by passing a current through it, the flame is coloured a deep green. To ascertain the diminution in the weight of a platinum wire when heated by the electric current, I placed between two clamping posts a wire five-thousandths of an inch in diameter, and weighing 266 m.grms. This wire, after it was brought to incandescence for twenty minutes by the current, lost 1 m.grm. The same wire was then raised to incandescence; for twenty minutes it gave a loss of 3 m.grms. Afterward it was kept incandescent for one hour and ten minutes, at which time it weighed 258 m.grms.—a total loss of 8 m.grms. Another wire, weighing 343 m.grms., was kept moderately incandescent for nine consecutive hours, after which it weighed 301 m.grms., showing a total loss of 42 m.grms. A platinum wire twenty-thousandths of an inch in diameter was wound in the form of a spiral one-eighth of an inch in diameter and one-half an inch in length. The two ends of the spiral were secured to clamping posts, and the whole apparatus was covered with a glass shade

* A Paper read before the American Association for the Advancement of Science; Saratoga Meeting.

2½ inches in diameter and 3 inches high. Upon bringing the spiral to incandescence for twenty minutes that part of the globe in line with the sides of the spiral became slightly darkened; in five hours the deposit became so thick that the incandescent spiral could not be seen through the deposit. This film, which was most perfect, consisted of platinum, and I have no doubt but that large plates of glass might be coated economically by placing them on each side of a large sheet of platinum, kept incandescent by the electric current. This loss in weight, together with the deposit upon the glass, presented a very serious obstacle to the use of metallic wires for giving light by incandescence, but this was easily surmounted after the cause was ascertained. I coated the wire forming the spiral with the oxide of magnesium, by dusting upon it finely powdered acetate of magnesium: while incandescent the salt was decomposed by the heat, and there remained a strongly adherent coating of the oxide. This spiral so coated was covered with a glass shade, and brought to incandescence for several minutes; but, instead of a deposit of platinum upon the glass, there was a deposit of the oxide of magnesia. From this and other experiments I became convinced that this effect was due to the washing action of the air upon the spiral; that the loss of weight in and the colouration of the hydrogen flame were also due to the wearing away of the surface of the platina to the attrition produced by the impact of the stream of gases upon the highly incandescent surface, and not to volatilisation, as commonly understood; and I venture to say, although I have not tried the experiment, that metallic sodium cannot be volatilised in high vacua by the heat derived from incandescent platinum; any effect that may be produced will be due to the washing action of the residual air. After the experiment last described I placed a spiral of platinum in the receiver of a common air-pump, and arranged it in such a manner that the current could pass through it, while the receiver was exhausted. At a pressure of 2 millimetres the spiral was kept at incandescence for two hours before the deposit was sufficient to become visible. In another experiment, at a higher exhaustion, it required five hours before a deposit became visible. In a sealed glass bulb, exhausted by a Sprengel pump to a point where a quarter of an inch spark from an induction-coil would not pass between points 1 millimetre apart, was placed a spiral, the connecting wires passing through the glass. This spiral has been kept at the most dazzling incandescence for hours without the slightest deposit becoming visible.

I will now describe other and far more important phenomena observed in my experiments. If a short length of platinum wire one-thousandth of an inch in diameter be held in the flame of a Bunsen burner, at some part it will fuse, and a piece of the wire will be bent at an angle by the action of the globule of melted platinum; in some cases there are several globules formed simultaneously, and the wire assumes a zigzag shape. With a wire four-thousandths of an inch in diameter this effect does not take place, as the temperature cannot be raised to equal that of the smaller wire, owing to the increased radiating surface and mass. After heating, if the wire be examined under a microscope, that part of the surface which has been incandescent will be found covered with innumerable cracks. If the wire be placed between clamping posts, and heated to incandescence for twenty minutes, by the passage of an electric current, the cracks will be so enlarged as to be seen with the naked eye; the wire, under the microscope, presents a shrunken appearance, and is full of deep cracks. If the current is continued for several hours these effects will so increase that the wire will fall to pieces. This disintegration has been noticed in platina long subjected to the action of a flame by Prof. John W. Draper. The failure of the process of lighting invented by the French chemist Tessie du Motay, who raised sheets of platinum to incandescence by introducing them into a hydrogen flame, was due to the rapid disintegration of the metal. I have ascertained the cause of

this phenomenon, and have succeeded in eliminating that which produces it, and in doing so have produced a metal in a state hitherto unknown, and which is absolutely stable at a temperature where nearly all substances melt or are consumed; a metal which, although originally soft and pliable, becomes as homogeneous as glass and as rigid as steel. When wound in the form of a spiral it is as springy and elastic when at the most dazzling incandescence as when cold, and cannot be annealed by any process now commonly known, for the cause of this shrinking and cracking of the wire is due entirely to the expansion of the air in the mechanical and physical pores of the platinum, and the contraction upon the escape of the air. Platinum as sold in commerce may be compared to sandstone, in which the whole is made of a great number of particles with many air spaces. The sandstone upon melting becomes homogeneous and no air spaces exist.

With platinum or any metal the air spaces may be eliminated, and the metal made homogeneous by a very simple process. This process I will now describe. I had made a large number of platinum spirals, all of the same size and from the same quality of wire; each spiral presented to the air a radiating surface of three-sixteenths of an inch; five of these were brought by the electric current up to the melting-point, the light was measured by a photometer, and the average light was equal to four standard candles for each spiral just at the melting-point. One of the same kind of spirals was placed in the receiver of an air-pump, and the air exhausted to 2 millimetres; a weak current was then passed through the wire, to slightly warm it for the purpose of assisting the passage of the air from the pores of the metal into the vacuum. The temperature of the wire was gradually augmented, at intervals of ten minutes, until it became red. The object of slowly increasing the temperature was to allow the air to pass out gradually and not explosively. Afterward the current was increased at intervals of fifteen minutes. Before each increase in the current the wire was allowed to cool, and the contraction and expansion at these high temperatures caused the wire to weld together at the points previously containing air. In one hour and forty minutes this spiral had reached such a temperature without melting that it was giving a light of twenty-five standard candles, whereas it would undoubtedly have melted before it gave a light of five candles had it not been put through the above process. Several more spirals were afterwards tried, with the same result. One spiral, which had been brought to these high temperatures more slowly, gave a light equal to thirty standard candles. In the open air this spiral gave nearly the same light, although it required more current to keep it at the same temperature. Upon examination of these spirals, which had passed through the vacuum process, by the aid of a microscope no cracks were visible; the wire had become as white as silver, and had a polish which could not be given it by any other means. The wire had a less diameter than before treatment, and it was exceedingly difficult to melt in the oxy-hydrogen flame. As compared with untreated platinum, it was found that it was as hard as the steel wire used in pianos, and that it could not be annealed at any temperature.

My experiments with many metals treated by this process have proved to my satisfaction, and I have no hesitation in stating that what is known as annealing of metals to make them soft and pliable is nothing more than the cracking of the metal. In every case where a hard drawn wire had been annealed a powerful microscope revealed myriads of cracks in the metal. Since the experiments of which I have just spoken I have, by the aid of Sprengel mercury pumps, produced higher exhaustion, and have, by consuming five hours in excluding the air from the wire and intermitting the current a great number of times, succeeded in obtaining a light of eight standard candles from a spiral of wire with a total radiating surface of 1-32nd of an inch, or a surface about equal to one grain of buckwheat. With spirals of this small size which have

not passed through the process, the average amount of light given out before melting is less than one standard candle. Thus I am enabled, by the increased capacity of platinum, to withstand high temperatures to employ small radiating surfaces, and thus reduce the energy required for candle light. I can now obtain eight separate jets, each giving out an absolutely steady light, and each equal to sixteen standard candles, or a total of 128 candles, by the expenditure of 30,000 foot-pounds of energy, or less than one-horse power. As a matter of curiosity I have made spirals of other metals, and excluded the air from them in the manner stated. Common iron wire may be made to give a light greater than platinum not heated. The iron becomes as hard as steel, and just as elastic. Nickel is far more refractory than iron. Steel wire used in pianos becomes decarbonised, but remains hard, and becomes the colour of silver. Aluminum melts only at a white heat.

In conclusion, it may be interesting to state that the melting-points of many oxides is dependent on the manner of applying the heat; for instance, pure oxide of zirconium does not fuse in the flame of the oxy-hydrogen blowpipe, while it melts like wax and conducts electricity when on an incandescent platinum spiral which is at a far lower temperature; on the other hand, oxide of aluminum easily melts in the oxy-hydrogen flame, while it only vitrifies on the platinum spiral.

A NEW METHOD OF PREPARING SULPHURETTED HYDROGEN.

By J. FLETCHER, F.C.S., London and Paris.

ANY mode by which the preparation of this useful gas can be rendered easier, and the unpleasantness of its manipulation diminished, will no doubt be welcomed by analysts: I therefore make no apology for submitting the results of some experiments made after reading a suggestion in some of the scientific journals, perhaps your own, but the name does not at the moment occur to me.

The plan is simply to fuse in a small glass flask sulphur and solid paraffin, leading the resulting gas by means of a perforated cork, india-rubber, and glass tube directly into the solution to be tested. The first gases are not sulphuretted, but when the mixture has been thoroughly fused and mixed the sulphuretted hydrogen passes over abundantly.

The advantage of the process is that the moment the flame of the lamp is removed the evolution of gas ceases, and the little apparatus can be laid aside without fear of creating offensive smells. When used again, the gas passes at once when sufficiently heated.

A washing bottle seems unnecessary. I passed the gas for an hour through such a bottle, and the water, although most strongly impregnated with the gas, was fairly clear and limpid, showing only the usual appearances.

There are a few precautions to be taken. The mixture is inclined to bump when strongly heated, but a few pieces of broken tobacco-pipe shank prevent that. Care must be taken that when the lamp is removed, and the gas ceases to pass, that none of the solution is sucked back into the bulb; it is very easily prevented. A very strong heat should not be applied, as then distillations would commence and the product condense in the tube.

I believe the process to be a simple, cleanly, and elegant substitute for the old methods, and particularly well suited for small and private laboratories. How it would work in large ones I would like to hear from those who are in a position to try it.

Blue Colour observed in the Scintillation of the Stars.—C. Montigny.—The blue colour is most rarely observed in dry seasons and conversely most abundantly in wet years.—*Les Mondes*.

EXPANSION OF SOLID AND FLUID BODIES.

By HERM. FR. WIEBE.

ON page 610, "Annual Report for 1878 of the German Chemical Society" (*Deutsche Chemische Gesellschaft*), I published some short remarks on the expansion-coefficient of solid bodies in its relation to the atomic volume. This relation, formerly unnoticed, is likely to throw some light upon the mechanical causes, still wrapped in such profound darkness, of the expansion of solids and liquids, and to disclose relations of fundamental significance.

(1.) The product of an atomic volume and the cubical expansion-coefficient represents the absolute expansion of an atom.

The mere contemplation of the latter already offers for solid elements interesting relations. Iron, cobalt, and nickel are of equal absolute expansion; the absolute expansions of other elements belonging to the same natural group are related to one another, according to simple proportions. Zinc and cadmium, for instance, show a proportion of 2:3; arsenic, antimony, and bismuth, 1:3:4.

Representing the absolute expansions as a function of atomic weight, they seem to present a curve similar to that of atomic volumes. The position of members of the natural families on that curve is an analogous one; maxima give sulphur, selenium, and tellurium, and most probably aluminium, gallium (?), indium, and thallium.

The absolute expansion of atoms manifests itself to be a periodical function of the atomic weight.

(2.) Expansion of a body represents that part of heat necessary to supersede its cohesive power. Suppose—

Density of a body to be	d ,
Its atomic weight	a ,
Its cubical expansion-coefficient	α ,
Its specific heat	c ,
Its boiling-point	s ,
and Its melting-point	σ ,

we have the following relation:—

$$\frac{d}{\frac{a \cdot \alpha}{d \cdot c (s - \sigma)}} = 2 \quad \dots \dots (1)^*$$

In words, the inverted value of absolute expansion of an atom stands in the proportion 2 to that quantity of heat necessary to bring equal volumes during constant atmospheric pressure from the melting- to the boiling-point. The equation (1) may still be simplified, when—density solving in the fraction—we have—

$$2 a \cdot \alpha \cdot c \cdot (s - \sigma) = 1 \quad \dots \dots (1a)$$

The subjoined table contains a compilation of expansion values of a few bodies in figures. (See next page.)

(3.)† In gaseous substances the molecules are so far separated as no longer to influence one another. Liquid and solid bodies consist of molecular groups, which evidently are formed by those especial powers, emanating from the molecules themselves. At the boiling-point, as well as at that of melting, all bodies possess an equal degree of cohesion. Therefore, if we multiply the absolute expansion of an atom with the temperatures of these fixed points (the temperatures having been augmented by the inverted expansion-coefficient previously), the result will be comparable figure-values, all of which are multiples of the expansion-coefficient.

For brevity's sake we give in the following table in the column immediately following the elements, the figures representing the absolute expansion of the atom; the second and third columns contain respectively the melting-point on the common scale, and in so-called absolute temperature; the fourth shows the product of the absolute expansion and temperature; and the fifth the expansion-

* See "Annual Report for 1878 of the German Chemical Society," p. 788.

† The following is an abstract of a paper, which will be published in the next number of the above-mentioned report.

Element.	<i>d</i>	<i>a</i>	<i>a</i>	<i>c</i>	<i>s</i>	$\frac{d}{\frac{a \cdot a}{c \cdot d(s-\sigma)}} = 2$
S ..	2.04	31.98	0.0002670	0.1710	447	$\frac{238.9}{116.3} = 2.05$
Se ..	4.60	78.00	0.0001696	0.0801	700	$\frac{358.5}{177.97} = 2.02$
P ..	2.30	30.96	0.0003556	0.1900	278	$\frac{208.9}{102.4} = 2.04$
Hg ..	13.596	199.8	0.0001882	0.0333	355.8	$\frac{361.6}{179.2} = 2.02$

coefficient and a constant—characteristic with each matter—expressed by the letter *m*.

Element.	I. Absolute Expansion of the Atom per 1 deg.	II. Melting-point in Common Scale.	III. Absolute Temperature so-called.	IV. Product III. I.	V. Expansion Coefficient.	<i>m</i> .
S	0.003015	113.6	388.6	1.171629	0.003905	300
Se	0.001872	217	492	0.921024	0.003607	250
Te	0.001029	489	764	0.786156	0.003931	200
Zn	0.000795	412	687	0.546165	0.003641	150
Cd	0.001188	315	590	0.700920	0.003505	200
As	0.000222	500	775	0.172050	0.003721	50
Sb	0.000630	430	705	0.444150	0.003701	120
Bi	0.000864	264	539	0.465696	0.003725	125
In	0.001911	176	451	0.861861	0.003591	240
Tl	0.001557	290	565	0.879705	0.003665	240
Pt	0.001530	334	609	0.931770	0.003728	250
Fe	0.000255	1600	1875	0.478125	0.003678	130
Co	0.000255	1500	1775	0.452625	0.003621	125
Ni	0.000255	1450	1725	0.439875	0.003665	120

Without at present enlarging upon these constants as to their explanation we simply wish to point out the fact that they seem to be in relation with the number of atoms, combining to form a solid molecular group; constants show, for elements of the same chemical group, simple proportions.

To the boiling-point similar proportions are applicable; the following relation, therefore, is generally adopted:—

$$\frac{a \cdot a}{d} \cdot T = \beta \cdot m \dots \dots (2)$$

in which β represents the expansion-coefficient in a gaseous matter; *T* the interval of temperature from the boiling- to the melting-point, or from the latter to the so-called absolute zero-point; and *a* the medium expansion-coefficient within these intervals of temperature.

It appears that the "Gay-Lussac rule" is generally valid, quantitatively changed only in its application to the fluid and solid state of matter; hence a common law of expansion is applicable to the whole scale of the state of aggregation in matter.

(4.) Applying the equation (2) to homologous rows of organic bodies, proportions astonishingly simple are the result.

The following table contains in column I. the medium absolute expansion of a molecule between melting- and boiling-points, or, the melting-point not being known, between 0° and the boiling-point. The figures of columns II. to V. answer to those in the preceding table.

Substance.	I. Medium Abs. Expansion of the Molecule per 1 deg.	II. Boiling-point in Common Scale.	III. Boiling-point from the III. I. Scale. Abs. Zero-point so-called.	IV. Product	V.
Formic acid ..	0.04326	100	375	15.6	5.2. 3
Acetic acid ..	0.06828	117.3	392.3	26.2	5.2. 5
Butyric acid ..	0.10235	144	421	46.8	5.2. 9
Methylic alcohol	0.05000	66.3	341.3	17.06	8.5. 2
Ethylic alcohol	0.07143	78.3	353.3	25.26	8.5. 3
Amylic alcohol	0.12500	131.8	406.8	50.8	8.5. 6

In this manner general formulæ for the boiling-point^s of both these homologous rows are easily established.

For the acids, the constant 5.2 is to be multiplied with the —by one augmented—number of atoms of hydrogen contained in gaseous molecule; for the row of alcohols, the constant 8.5 with half the number of atoms of hydrogen.

We purpose treating the above subject more in *extenso* very soon, especially to enlarge upon the theoretic significance of the constant *m*.

Berlin, September, 1879.

CORRESPONDENCE.

DISSOCIATION OF CHLORINE.

To the Editor of the Chemical News.

SIR,—Mr. F. P. Dunnington, in his short notice on the "Dissociation of Chlorine, seems to have mistaken PtCl_4 for Pt_2Cl_4 (PtCl_2), two very different chlorides. If he refer to the quoted article, CHEMICAL NEWS, vol. xl., p. 69, or to p. 49, he will find PtCl_2 is the chloride Victor Meyer used. Due care was taken to exclude the possibility of oxygen compounds being present, and more, extraordinary care was taken. The PtCl_2 (Pt_2Cl_4) was amply proved to be chemically pure before use, a point which it can hardly be thought for an instant a chemist of European reputation like Victor Meyer would overlook or neglect. It is very evident that Mr. Dunnington had succeeded better in preparing his own pure chlorine if he had but used this very PtCl_2 (Pt_2Cl_4) for the purpose, as Victor Meyer did. —I am, &c.,

WATSON SMITH, F.C.S., F.I.C.

City of London College.—Evening Classes for Young Men, 52, Leadenhall Street, E.C.—The Prospectus of this Institution has been published, stating that the new Session commences on Monday, October 6th. The Inaugural Address will be given by the Right Rev. the Bishop of Bedford, Bishop Suffragan of London, on Thursday evening, October 9th, at 8 o'clock, the subject being "Books, and how to use them." A glance at the Prospectus shows that the Council of the College are extending its curriculum, the array of classes being greater than in any previous session. The need of a much larger building has been thoroughly proved, and it is hoped that the exertions of the Council to obtain a site for a new college will soon be successful. It surely is not too much to expect that some of the wealthy citizens of this great metropolis, emulating the example set by many in several provincial towns, will come forward with a portion of the fortune which they have amassed, in aid of so worthy an object as the rearing of a fine and commodious building for the express purpose of improving, mentally and morally, the young men on whom these merchant princes depend for the proper conduct of their business. No more philanthropic object can engage their attention.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 8, August 25, 1879.

Digestive Ferment of Carica Papaya.—A. Wurtz and E. Bouchut.—From the undecomposed juice of this tree the authors have obtained by precipitation with alcohol, the ferment, in the form of an amorphous white powder, entirely soluble in water, and containing 10 per cent of nitrogen. To this substance they have provisionally given the name of papain. It is distinguished from pepsin by the circumstance that it is capable of dissolving large quantities of fibrin not merely in presence of a small quantity of acid, but even in a neutral or slightly alkaline medium. It is doubtless analogous to the ferments secreted by carnivorous plants, such as *Nepenthes*, *Drosera*, &c.

Reply to the Observations of M. Berthelot.—A. Wurtz.—A continuation of the chloral controversy.

Compressibility of Gases at High Pressures.—E. H. Amagat.—It appears very probable that when a gas submitted to increasing pressures, whether or not it has shown at first an augmentation in its compressibility, presents afterwards a decrease, it is always placed in conditions where it may, by pressure alone, pass gradually through all the intermediate conditions between the gaseous and the liquid state without liquefaction properly so-called. The decrease in the compressibility indicates then in general that the gas has reached a temperature higher than that of the critical point.

Maximum Tension and the Density of the Vapour of Alizarin.—L. Troost.—The maximum tension of the vapour of alizarin is about 11 m.m. at 261°, and 20 m.m. at 276°. The density observed is 16.32 as against 16.62 by calculation.

Purification of Hydrogen.—A. Lionet.—Oxide of copper in the cold arrests all the combinations of hydrogen which may be present as impurities, except the hydrogen carbides.

Active Principle of Ammi Visnaga.—M. Ibrahim Mustapha.—The author has isolated a crystalline matter, which behaves like a glucoside. The composition of this principle, for which the author proposes the name kellin, is not given.

Distribution of Copper in the Primitive Rocks and in their Derived Sedimentary Deposits.—L. Dieulefait.—Copper is disseminated throughout the entire thickness of the primordial formation. In no case is it requisite to operate on more than 100 grms. of rock in order to isolate the metal. It is also found in the ordinary sedimentary deposits, such as the silurian and infra-silurian of Algeria, &c. The last mother-liquors of the salt marshes of the Mediterranean contain copper in such quantity that it may be easily detected in 5 c.c.

No. 9, September 1, 1879.

Note on the Solar Temperatures.—J. Jansen.—The author considers that the nucleus of the sun is much hotter than the photosphere or any of the exterior strata. The expression "solar temperature" is deficient in precision. The calorific and thermo-electric methods generally employed seem capable of a rational use for determining the thermic power of the solar radiation which arrives at the earth's surface, but they cannot give exact notions as to the real temperatures of the sun, not even for its mean temperature, an expression which as applied to the sun seems almost without meaning.

The Chemical Constitution of Alkaline Amalgams.—M. Berthelot.—The author considers these amalgams as of great importance as the type of compounds resulting from the union of two solid constituents, such as the metallic alloys, the cryohydrates, the fats, butters, resins, &c. Are these products formed by the simple mixture of certain definite compounds, associated sometimes with each other, sometimes with one of the components in excess, in the manner of two powders mechanically mixed, and then brought into a coherent mass by external pressure? Or are the properties of each of these definite compounds modified more profoundly by the presence of an increasing dose of the other definite compound or by that of the component in excess, so that the properties of the total mass cannot be represented by the pure and simple sum of those of the two definite bodies supposed to be mixed? These questions the author endeavours to solve by thermo-chemical methods.

Partial Synthesis of Milk-sugar, and Contribution to the Synthesis of Cane-sugar.—E. Demole.—The author recognises a profound difference between milk-sugar and cane-sugar. Two mols. dextro-glucose, reunited and with loss of water, will not in any case reconstitute cane-sugar. Milk-sugar, in contact with dilute acids, is resolved into two isomers, galactose and lacto-glucose. These products, freed from acid and carefully evaporated to dryness, were treated with 3 parts of acetic anhydride, yielding a body having all the properties of octacetylated milk-sugar.

Reaction of Tungstates in Presence of Mannite.—M. Klein.—There is a parallelism between the reactions of the biborates and those of the para-tungstates and meta-tungstates, which extends even to the properties of the compounds which these bodies form with mannite, compounds otherwise devoid of interest.

Determination of Urea; a Reply to M. G. Esbach (*Comptes Rendus*, Aug. 18, 1879).—C. Méhu.—The author states that in presence of pure sugar sodium hypobromite liberates all the nitrogen of the urea and nothing more.

No. 10, September 8, 1879.

Compounds of the Hydracids with Ammonia.—E. J. Maumené.—If crystals of ammonium hydrosulphate reduced to a fine powder are added to very concentrated ammonia, cooled down to 0°, we obtain after the lapse of some hours transparent crystals of the mean composition $\text{HS}(\text{H}_3\text{N})_2$, that is to say, a very basic hydrosulphate. Theory enables the author to predict the existence of two series of four members each, in one of which the ammonia predominates, and in the other the sulphuretted hydrogen.

Any two compounds, either of one or both of these series may again combine among themselves so that the compounds of HS and H_3N are innumerable. The compounds of hydrochloric acid and ammonia offer analogous relations.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 18, 1879.

Action of the Aldehyds upon Chloral-ammonia.—R. Schiff.—By mixing equivalents of dry chloral-ammonia and benzaldehyd, a compound was obtained of the composition $\text{C}_9\text{H}_8\text{Cl}_3\text{NO}$. Valeraldehyd, furfurol, cinnanhol, and acetaldehyd exert a similar action.

On Penta-brom-resorcin.—R. Benedikt.—This product, which Liebermann and Dittler have felt it incumbent upon them to re-name tribrom-resoquinon bromide, has been examined as to its behaviour with reducing agents, when tri-brom-resorcin is produced, and with aniline and phenol.

A Hydrocarbon obtained from Rosin Oil by Heating with Sulphur.—W. Kelbe.—If the highest-boiling products of the dry distillation of rosin are heated with sulphur to 200°, a crystalline body is obtained consisting of 91.5 per cent carbon and 8.5 hydrogen.

Derivatives of Oxy-propyl-benzoic Acid.—R. Meyer and J. Rosicki.—An examination of para-diphenyl-benzol and of its further derivatives.

A New Reaction of Kreatinin and Kreatin.—Th. Weyl.—If a few c.c. of recent human urine are mixed in a test-tube with a few drops of a very dilute solution of sodium nitro-prusside, and if dilute soda-lye is then added drop by drop, a beautiful ruby-red colour appears, which in a few minutes changes to an intense straw-yellow. This reaction seems characteristic of kreatinin. As kreatin is converted into kreatinin by boiling with dilute sulphuric acid, the reaction may serve also for the detection of this substance.

Aceto-propionic Acid, and its Identity with Levulinic Acid.—Max Conrad.—The melting-points, boiling-points, specific gravities, indices of refraction, and powers of dispersion of the two acids are identical.

A Correction.—P. Griess.—The author points out an error in the formula of amido-benzoic-percyanide as given in his paper (*Berichte*, xi., 1875).

Preliminary Communication.—G. Vortmann.—This notice relates to the cobalt-ammonia compounds.

Perchloro-phenol-chloride.—F. Beilstein.—This substance was repeatedly formed on passing chloride into an acetic solution of the acetyl compound of meta-chloranilin.

Accuracy of Thermo-chemical Numerical Results.—Julius Thomsen.—The author replies to M. Berthelot by pointing out a number of inaccuracies in the results published by the latter.

Action of Hypochlorous Acid upon Ethylen.—L. Pebal.—A preliminary communication, in which the author's own results do not appear.

Aspido Spermin, an Alkaloid of the Quebracho Bark.—G. Fraude.—Aspido spermin crystallises in small white prisms with strongly shining surfaces, readily soluble in alcohol and ether, but dissolving sparingly in water, and melting at 205° to 206°. The taste of the solutions resembles that of quinine.

New Investigations on the Diazo-compounds.—P. Griess (Part vi.).—In this valuable paper, which unfortunately does not admit of abstraction, the author examines the action of certain sulpho-diazo-acids upon the phenols.

New Glucoside from Lupinus luteus.—E. Schulze and J. Barbieri.—The substance obtained is lupinin, which is readily split up into lupigenin and a glucose, probably dextrose.

Transformation of Acetaldehyd into Mercaptan.—C. Böttinger.—Not adapted for abstraction.

Hydrazin Compounds of the Fatty Series.—E. Fischer.—The author describes the preparation of diethyl-hydrazin, its oxidation, and its behaviour with nitrous acid.

Contributions to the Voluminar and Steric Laws.—H. Schröder.—In this memoir the author applies his theory to the compounds of thallium and ammonium.

Dimethyl-acrylic Acid.—W. von Miller.—The author obtains this acid by distilling with sulphuric acid an oxidation product arising from the action of permanganate upon isobutyl-formic acid.

Actions of Nitrosyl Silver.—W. Zorn.—An examination of the behaviour of nitrosyl silver with ammonium chloride, with methylamin and aniline hydrochlorates, phenol and acetester.

Transformation of Undecylenic Acid into Undecylic Acid.—F. Kraft.—Not suitable for abstraction.

Absorptive Power of Arable Earth, and of Silicic Acid.—J. M. v. Bemmelen.—In opposition to Liebig, the author refers this absorption not to physical surface attraction, but to chemical action.

On Propyl-benzoic Acid.—E. Paterno.—The author points out that propyl-benzoic acid, described as new by

H. Korner (*Berichte*, p. 1863), was described by Spica and himself in June, 1877.

Action of Chloro-carbonic Ethylether upon certain Oxygenated Haloid Compounds of the Fatty Series.—O. J. Kelly.—The author examines the behaviour of chloro-carbonic ethylether with dibrom-allylic alcohol, with dichlorhydrin, and epichlorhydrin.

Contribution to the Knowledge of Alstonia Bark.—O. Hesse.—The alstonin of palm is a mixture of chlorogenin and porphyrin, and the alstonin of F. v. Müller and Rummel an impure chlorogenin.

The Initial Members of the Paraffin Series on Exhaustive Bromation.—V. Merz and W. Weith.

The Final Products of the Exhaustive Bromation of certain Paraffins having Higher Molecules.—V. Merz and W. Weith.—Methan at 180° is converted by direct bromation into perbrom methan. Carbon sulphide, under the influence of iodiferous bromine, undergoes this metamorphosis, partially at least at common temperatures. On heating perbrom-methan the immediate product is perbrom-ethylen, and at 350° perbrom-benzol. Ethan, ethyl iodide, and ethylen dibromide, in contact with iodiferous bromine, are converted into perbromised ethylen and ethan at 200° to 250°. The hydrocarbons of the fatty series behave on energetic bromation exactly as they do on energetic chloration, save that bromine reacts less readily than chlorine.

Process for the Determination of Vapour-densities of Bodies boiling above 440°, as also of such Bodies as act upon Mercury and Wood's Metal.—V. Meyer and C. Meyer.

On Substituted Phthalinites.—S. Gabriel.—A description of chlorphenyl-phthalimid, of brom- and iod-phenyl-phthalimid, of nitro-phenyl-phthalimid, and of ortho- and meta-phthalimido-benzoic acid.

Certain Derivatives of Ortho-nitro-phenol.—Jos. Bendix.—The author attempting to produce dioxy-diphenyl-sulphurea, obtained instead oxy-phenyl-sulphurea.

On Orthotheoformic Benzyl-ether.—M. Dennstedt.—The author has obtained and analysed this body, which has the composition $C_{22}H_{22}S_3$.

Action of Sulphuric Anhydride upon Pseudo-sulpho-cyanate of Phenyl.—G. Magatti.—Noticed under *Gazzetta Chimica Italiana*.

Synthesis of Oxyketons by the Introduction of Acid Radicles into Phenols.—O. Doebner and W. Stackmann.—In this communication the authors describe the formation of oxyketons by the action of benzoyl chloride upon resorcin.

On Malachite Green.—O. Doebner.—A controversial paper directed against E. and O. Fischer. The author maintains that the composition of malachite green is $C_{23}H_{24}N_2$; that its reduction-product, $C_{23}H_{26}N_2$, is identical with the base obtained from the oil of bitter almonds; and that the green colouring-matters formed on the oxidation of this base are not proved to be identical with malachite green.

Methylated Anilines and Toluydins, and their Coloured Derivatives.—P. Monnet, F. Reverdin, and E. Noelting.—Noticed elsewhere.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 16, August 14, 1879.

Prof. Sonnenschein has given the somewhat singular name of "Virginia" to a substance which he has extracted from the residues of the distillation of petroleum. He describes it as a semi-transparent mass, yellowish, and fatty, and displaying, when heated, a blue fluorescence. It melts at 47°, and is partially soluble in ether.

No. 17, August 21, 1879.

The French Association for the Advancement of Science has held its annual meeting at Montpellier from Aug. 24 to Sept. 4, under the presidency of M. Bardoux. Public lectures have been delivered on the Rhone irrigation canal and on the electric light.

In noticing the proceedings of the International Office of Weights and Measures, M. Moigno suggests that it would be preferable to adopt as the true unit of measure the cubit of the great Pyramid, the same as the inspired cubit of Moses and Solomon, the ten-millionth part of the polar axis of the globe. He proposes that the meridian of the pyramid should be adopted by all nations as the first meridian.

M. l'Abbé le Dantec has constructed an electric machine which doubles the force of the Carré machine. He obtains from it continuous sparks of 15 centimetres.

Les Mondes will be carried on for the future by a limited liability company, the editorship remaining unchanged.

No. 18, August 28, 1879.

A New "Splendeur de la Foi."—M. Moigno proposes an exploring expedition to the Red Sea and the isthmus of Suez for the purpose of discovering relics of the army of Pharaoh. A branch-expedition is to seek for the aërolites which fell from heaven in the famous battle gained by Joshua, and which covered the soil about Bethoron. M. Moigno suggests that the sale of these stones would to a large extent cover the expense of the expedition. He takes occasion to misconstrue the language used by Prof. Richard Owen concerning the isthmus of Suez at the Congress of Orientalists, held in London in September, 1874, accusing him of making the Red Sea disappear from the region traversed by the Israelites.

No. 1, September 4, 1879.

This issue contains no original matter in physics or chemistry. Two articles are devoted to attacks upon Prof. Paul Bert, in the latter of which the editor states that he for the first time inserts a paper on religious polemics!

No. 2, September 11, 1879.

This issue again contains no original chemical or physical matter. There is an article on the *Splendeurs de la Foi*, and a very violent attack upon the French Government for wishing to keep in its own hands the right of granting Academical degrees.

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NOTE ON THE
BEHAVIOUR OF DOUBLE CHLORIDE OF
COPPER AND AMMONIUM
WITH FERROUS SULPHIDE.

By W. F. K. STOCK, F.C.S., F.I.C.

WHILST experimenting recently on the accurate determination of carbon in iron or steel containing much sulphur, it appeared to be desirable to ascertain definitely in what manner the reagent used for the carbon separation operated upon the iron-sulphur compound, but as the actual composition of that compound could only be speculated upon in the vaguest light, if at all, it was thought best to work upon a sulphide of known quality, and afterwards, if the preliminary experiment showed it to be feasible, to endeavour to separate and examine the unknown sulphide.

The process used for the carbon separation was that known as McCreath's method, and consisted in treating a weighed quantity of iron or steel with a hot concentrated solution of double chloride of copper and ammonium ($2\text{NH}_4\text{Cl}, \text{CuCl}_2, 2\text{H}_2\text{O}$), the reaction depending upon the simultaneous formation of ferrous and cuprous chlorides, the carbon being left in a free state.

A tolerably pure ferrous sulphide was prepared by heating a bar of good wrought-iron to whiteness in a coke fire, and plunging it into a Cornish pot containing melted sulphur, and when the greater part of the sulphur was used up the pot was closely covered and heated to bright redness, the excess of sulphur being thus got rid of.

A careful analysis of the pounded and mixed sulphide gave the following figures:—

Iron.. ..	63.98 per cent
Sulphur	33.86 "
Sand, &c.	2.16 "
	100.00 "

Theory requires for FeS —

Iron.. ..	63.63 per cent
Sulphur	36.37 "
	100.00 "

It will be seen that in the prepared sulphide the iron is very slightly in excess of the sulphur.

On treating 0.5 grm. of the powdered sulphide with 50 c.c. of the concentrated copper-ammonium solution at nearly boiling heat for half an hour, the original blackish grey colour of the sulphide was changed to yellowish grey, and upon filtering off the residue, washing first with boiling water, next with strong alcohol, and finally exhausting with carbonic disulphide, a great part of the residue was found to have been dissolved by the latter menstruum, which, upon spontaneous evaporation, yielded a crop of micro-crystals, and these under a one-inch objective, and viewed with binocular vision, afforded some exquisitely perfect rhombo-octahedra.

It thus became evident that the action of the double copper-ammonium salt upon iron-carbide and iron-sulphide was precisely analogous, and that the method held out no hope of separation. It only remained to find to what extent the decomposition had proceeded during the half hour's exposure, and this was effected by igniting the residue, after treatment with carbonic disulphide, in a

muffle with free access of air, and weighing as Fe_2O_3 . 0.5 grm. sulphide gave 0.0954 residual Fe_2O_3 , which is equal to 19.09 per cent undecomposed sulphide. Hence, allowing for oxidation during washing, &c., it may safely be assumed that 80 per cent of the original sulphide was decomposed by the double copper-ammonium salt, with a proportional liberation of free sulphur.

A second experiment was made with native ferric sulphide, which was very finely powdered and exposed at boiling heat for over an hour to the copper solution as before, but, although some free sulphur was obtained, the decomposition was far from complete.

ON SCANDIUM.

By M. P. CLÈVE.

In the sitting of March 24, 1879, M. Nilson announced the discovery of a new element, which he named scandium, and which he had extracted from ytterbia. Some weeks after this publication I found the same metal in the gadolinite and yttrötitanite of Norway, and have examined its characters.

It occurs in minute quantities only. Gadolinite contains 0.002 per cent of the oxide, and yttrötitanite 0.005. Its only oxide, scandia, has the formula Sc_2O_3 . The composition of the double ammoniacal sulphate and the double potassic oxalate proves the correctness of this formula. The atomic weight of scandium = 45.12. Scandium oxide, or scandia (Sc_2O_3), is a perfectly white, light, infusible powder resembling magnesia. Even the strongest acids attack it with difficulty, yet it is more soluble in acids than is alumina. Sulphuric acid yields with scandia sulphate as a white voluminous mass resembling thorium sulphate in appearance. Scandia dissolves more readily in hydrochloric than in nitric acid. The specific gravity of the oxide is approximately 3.8. The hydrate is a white bulky precipitate, resembling hydrate of alumina. If dried it forms semitransparent fragments. It does not appear to absorb the carbonic acid of the air. It is insoluble in an excess of ammonia or of caustic potassa. It does not decompose sal-ammoniac if heated together.

The salts of scandium are colourless or white; they possess a very harsh, astringent taste, quite distinct from the sugary flavour of the other yttria earths. The sulphate does not form distinct crystals; the nitrate, formate, oxalate, and acetate are crystallisable. The chloride gives the following reactions:—If heated in the gas-flame it does not produce a spectrum. Potassa and ammonia give a voluminous precipitate, insoluble in excess of the reagents. Tartaric acid prevents precipitation in the cold, but if the solution is heated an abundant and voluminous precipitate appears. Sodium carbonate produces a voluminous precipitate, soluble in excess. Sulphuretted hydrogen effects no change. Ammonium sulphide precipitates the hydrate. Sodium phosphate gives a gelatinous precipitate. Oxalic acid produces a curdy precipitate, which is speedily transformed into a crystalline powder. The oxalate is soluble in concentrated acids, and its separation from an acid solution is not complete. Although the oxalate appears more soluble than the oxalates of the other earths it is found in the first fractions when a mixture of salts of scandium and ytterbium is submitted to partial precipitation. The acid oxalate of potassa gives a double salt in a crystalline powder. Sodium hyposulphite precipitates a boiling solution readily, but not completely. Sodium acetate gives a precipitate at the boiling-point, which is readily detached from the filter, but the precipitation is incomplete. In concentrated solutions potassium sulphate occasions the separation of a double salt as a crystalline powder, soluble in a saturated solution of potassium salt. Sodium sulphate behaves in the same manner. The author has described in his memoir the chloride, nitrate, sulphate, three double sul-

phates, a double oxalate, the selenite, acetate, and formiate. The great interest of scandium is that its existence had been predicted. Mendeleef, in his memoir on the law of periodicity, had foreseen the existence of a metal which he named ekabor, and whose characters agree very fairly with those of scandium.

Predicted Characters of Ekabor.

Atomic weight = 44.
Ekabor should have only one stable oxide, Eb_2O_3 , a base more energetic than alumina, with which it has several characters in common. It should be less basic than magnesia.

Though yttria should be a more energetic base we may foresee a great resemblance between yttria and oxide of ekabor. If ekabor is found mixed with yttrium their separation will be difficult and will be founded on differences of solubility and of basic energy.

Oxide of ekabor is insoluble in alkalies; it is doubtful if it will decompose ammonium chloride.

Its salts will be colourless and give gelatinous precipitates with KOH , Na_2CO_3 , and HNa_2SO_4 .

With potassium sulphate it will form a double salt having the composition of alum, but scarcely isomorphous with that salt.

Few of the salts of ekabor will crystallise well.

Water will decompose the anhydrous chloride of ekabor with liberation of hydrochloric acid.

The oxide will be infusible, and after ignition will dissolve in acids with difficulty.

The specific gravity of the oxide will be about 3.5.

—*Comptes Rendus.*

Observed Characters of Scandium.

Atomic weight = 45.
Scandium has only one oxide, Sc_2O_3 , a base much more energetic than alumina, but feebler than magnesia.

Scandia is less basic than yttria, and their separation depends on the different stability of their nitrates on exposure to heat.

Hydrate of scandium is insoluble in alkalies and does not decompose ammonium chloride.

Salts of scandium are colourless and give gelatinous precipitates with KOH , Na_2CO_3 , and HNa_2SO_4 .

The double sulphate of ammonium and scandium is anhydrous, but has in other respects the composition of alum.

Scandium sulphate does not form distinct crystals though the nitrate, acetate, and formiate are crystalline.

The crystalline chloride is decomposed and evolves hydrochloric acid when heated.

The ignited oxide is an infusible powder and dissolves in acids with difficulty.

The specific gravity of the oxide is 3.8.

Mr. Narjes, an engineer connected with Herr Krupp's works at Essen, who, on the 16th and 17th March, 1877, worked the first heat on a large scale, 4 tons of pig, holding 0.7 per cent. of phosphorus, being reduced to metal running 0.134 per cent., while the percentage of carbon sunk only from 3.1 to 3.03 per cent. Narjes' process consists in dephosphorising and refining the pig, without affecting the carbon percentage materially, by oxides of iron and manganese, partly used as fettling and partly as additions. The practice at Essen is said to be simple; the pig is melted in a 13-foot cupola with coke in one hour and a half, and is then tapped into a furnace similar to the Pernot, heated by the regenerative system. The flat hearth is covered with a layer of almost one foot of iron ore melted on at a very high temperature. Before every heat from 1500 to 1700 pounds of ore, also heated until sintered, are added before the iron is tapped from the cupola. The temperature during the operation of dephosphorising is a strong white heat, which constitutes the difference between the Krupp and the Bell methods. At first the furnace is made to revolve twice in a minute, which is soon increased to five. Generally after five, and never after more than ten minutes, the dephosphorisation and refining are completed—a point which is indicated by the commencement of bubbling and the appearance of small flames of carbonic oxide. The following analyses, made by Professor Tinkener, of the Berlin School of Mines, will show that dephosphorisation is effected without any diminution of the percentage of carbon, 'a' designating before and 'b' after the process:—

	I a.	I b.	II a.	II b.	III a.	III b.
" Carbon	{ 3.39 3.98	{ 3.75 3.77	{ 3.81 3.78	{ 3.56 3.57	{ 3.17 3.16	{ 3.02 3.04
" Phosphorus	{ 0.632 0.629	{ 0.131 0.133	{ 0.45 0.445	{ 0.108 0.106	{ 1.223 1.218	{ 0.303 0.301

"All the analyses, it will be seen, were made in duplicate. It is expected, although it has not yet been proved experimentally, that manganiferous pig will work more favourably still. Another point which has not yet been settled is whether it will be possible, by adding a silicious pig, to fit the refined metal for the Bessemer process, for which, as at present constituted, it is not suitable, as the dephosphorising process eliminates the silicon simultaneously."

Dr. J. M. Brown pointed out recently that phosphorus and silicon were largely removed, and carbon completely, by heating with sodium or other alkaline carbonates in a platinum vessel.

Herr Krupp, of Essen, is said to have forced through a bath of molten iron streams of fused alkaline chlorides, fluorides, &c., and with some measure of success.

The late Mr. William Baker, of this town, found that a distinct quantity of phosphorus was removed by blowing chlorine through molten iron in a plumbago crucible, but that in an ordinary fire-clay crucible the action was but slight. He attributed some of the success of this experiment to the presence of an excess of carbon.

Almost every salt known to chemists has been tried from time to time, but most frequently by people who could give no reason for their suggestions. The preceding may, perhaps, be taken as the most noteworthy processes possessed of any reasonable symptoms of practical success which were known up to the present, with the single exception of Mr. Snelus's experiment in 1872.

Mr. G. J. Snelus in that year found that phosphorus could be eliminated in the Bessemer converter by forming a basic slag, and he effected this by lining the vessel with basic material, in contrast to the usual gannister or acid lining. Silica being a more powerful acid than phosphoric displaces the latter from combination at high temperatures; consequently, unless the silica is thoroughly saturated, there can be no free base for the phosphoric acid to combine with, and therefore, in contact with iron at high temperatures, it will be reduced as fast as made, if, indeed,

ON THE
SEPARATION OF PHOSPHORUS AND IRON,
ESPECIALLY WITH REFERENCE
TO THE MANUFACTURE OF STEEL.*

By THOMAS BLAIR.

(Concluded from page 152.)

Krupp's Process for Dephosphorising Pig-Iron.—Professor Wedding gives some data on the practical working of Krupp's, or rather Narjes', process for dephosphorising pig-iron. "The originator of the process is

* Read before the British Association for the Advancement of Science Section B., Sheffield, 1879.

the phosphorus is oxidised at all under these conditions. These considerations led Mr. Snelus to believe that phosphorus was removed in all processes, just in proportion to the basic nature of the slag.

Observing that certain kinds of limestone, the magnesian variety especially, when submitted to very intense heat, became indurated, and converted into a form in which they were no longer acted on by water, Mr. Snelus lined up a 2 cwt. converter with lime made from this stone, and fired off at a very high temperature. By keeping the slag basic he successfully dephosphorised Cleveland pig-iron. His results were eminently satisfactory, proving beyond a doubt that he had hit upon the proper method. Unfortunately, Mr. Snelus allowed his process to lie dormant, and was thus shorn of the glory which would otherwise have been his. His statement made at the Iron and Steel Institute in May this year was conclusive though tardy.

Messrs. Thomas and Gilchrist have, with great energy and perseverance, been able to succeed chemically in producing good steel from Cleveland pig. Their first experiments were made at Blaenavon, in South Wales, and their results there were sufficiently satisfactory to induce practical steel-makers to take up the subject. Mr. E. Windsor Richards, of Messrs. Bolckow, Vaughan, and Co.'s works, Middlesbrough, went into the matter so thoroughly that steel rails have already been made in considerable quantities by this process.

Briefly, the process of Messrs. Thomas and Gilchrist consists of the following points:—

1. A durable basic lining.
2. The addition of basic materials.
3. Removal of phosphorus by blowing after the carbon has been eliminated.

In the first place immense difficulties have been experienced in producing a durable basic lining. Caustic lime, as is well known, is not stable under ordinary atmospheric conditions. But, as has already been pointed out, Mr. Snelus was able to select certain kinds of magnesian limestone which became indurated by calcination at very high temperatures. Linings have thus been made, which seem tolerably permanent. A brick and some pieces are on the table, and have the following composition:—

Silica	9·2 per cent.
Lime	46·7 "
Magnesia	32·8 "
Oxide of iron, alumina, alkalies, &c.	11·7 "

It will thus be seen that a very successful result has been arrived at, as this brick is extremely hard and appears durable in the air, since it has been kept for some months without protection.

The method recently adopted was simply to form the mould as usual with such a mixture of the various seams of the magnesian limestone as would give a composition similar to the analysis given; to dry the brick gently; fire it in a kiln—first, moderately; then at a most intense white heat, so intense, indeed, that ordinary silica bricks are melted. Silicate of soda has also been used as a binding material.

Mr. Riley has found that quick-lime may be rendered plastic by admixture with petroleum or kindred oils, and either moulded into bricks or rammed in the vessel as is done with ordinary gannister.

The writer is informed that, in practice, linings made by either of these processes, when using irons not containing an excess of silicon, answer extremely well.

To prevent the oxidised silicon from attacking the lime lining, additions of lime (with or without admixture with oxide of iron) are made. This proceeding is a very wise one, and is found to answer well. Similarly in the reverse case, in the Hollway metallic sulphides process, in which large quantities of ferrous oxide are formed by oxidation, in a gannister or silicious-lined vessel, additions of powdered gannister are found to be very effective in miti-

gating the corrosive action of the oxide on the lining. Samples of the various materials used for refractory linings are shown.

2nd. The vessel being lined with the basic bricks and ready for use, about 6 tons of Cleveland pig-iron are poured in, and with or before this a quantity of about a ton of the basic material. This same vessel would take 8 tons of ordinary hæmatite Bessemer pig. The blow then commences, and differs but little from the ordinary Bessemer blow until the carbon is burnt out.

3rd. At this moment the vessel is turned down, and a sample is taken which is rapidly hammered out and broken, and from the appearance of the fracture and its brittleness a rough estimate is formed of the amount of phosphorus remaining. The vessel is then turned up, and the charge is blown for about thirty seconds, and a second sample is taken and examined as before. This operation is repeated, with diminishing times, until a sample is obtained which shows a small fine grain, and is sound on the edges. From 8½ to 10 per cent of spiegeleisen is then added, as in the ordinary process, but in this case a somewhat violent reaction takes place between the metal and spiegel, owing most probably to the fact that some amount of oxide of iron is formed, which is rapidly attacked by metallic manganese. The metal is then poured into the moulds as usual, and sent to the mills.

The following are analyses of a blow by this process:—

	Pig-iron.	1st Samp.	2nd Samp.	3rd Samp.	Steel.
Graphite	2·74	—	—	—	—
Com. carbon	0·30	trace	trace	trace	0·150
Silicon	2·28	trace	—	—	0·018
Sulphur	0·07	0·074	0·068	0·063	0·060
Phosphorus	1·48	0·904	0·238	0·076	0·177
Manganese	0·64	0·070	0·070	0·050	0·500

It will be here noted that the silicon disappears rapidly, and that in three minutes the carbon begins to go also, and that but little of the phosphorus is oxidised until most of the carbon has gone, and all the silicon.

In the refinery, puddling, and Bell's processes, however, the phosphorus commences to leave the iron before all the silicon has gone, and the carbon goes at a comparatively slow rate. This difference may be accounted for by the very much lower temperature of these processes, and the presence of much oxide of iron. But it has been asserted by M. Pourcel, whose opinion is worthy of the highest respect, "that in the old refinery process the phosphorus disappears in direct proportion with the silicon. Thus, when a pig rich in silicon and phosphorus is refined, the phosphorus continues to pass into the slag as long as the silicon is burning, that is to say, as long as the carbon does not burn, and no carbonic oxide is formed by intermolecular action. If the content of silicon is high with respect to the phosphorus and carbon; if the medium in which the process is conducted furnishes no free silica, but, on the contrary, an excess of oxide of iron to neutralise the silica coming from the silicon in the pig, we succeed in scorifying the greater part of the phosphorus."

In the Krupp and Bell processes the period of dephosphorisation is determined by the first appearance of the blue flames of carbonic oxide, showing that the silicon is all oxidised, and that the carbon commences to burn. No further removal of phosphorus takes place, but inasmuch as a reducing action has succeeded to an oxidising one, the phosphorus returns to the iron.

M. Pourcel concludes from this that a highly silicious iron would be advantageous for use in the Thomas and Gilchrist process, but the writer would submit that he is judging from a wrong standpoint, inasmuch as that the carbon in this case is rapidly disappearing at the same time as the phosphorus, and that it is necessary to continue the blow after the carbon has gone solely to scorify the phosphide of iron.

So far, however, from the presence of silicon being an

advantage, the reverse is the case, since it not only prolongs the blow, but the silica formed by its oxidation exercises a corrosive action on the lime lining of the converter. Its only advantage is, that it generates a great amount of heat very rapidly during its combustion. Were it possible to obtain the same heat from other sources, it would be possible and profitable to use iron very low in silicon or as free as possible from it.

Various methods have been proposed to effect this, such as, for instance, the substitution of manganese and the addition of carbon, in some form, to the molten metal. Mr. Lowthian Bell, some time ago, proposed to blow in powdered charcoal with the blast. Mr. Hollway proposed to form a neutral gas by blowing air through incandescent coke, and then to introduce with this gas carbon or liquid hydrocarbon. Mr. Wilkes, of Manchester, has made some very promising experiments by simply running petroleum spirit into the ordinary blast pipes.

Mr. Hollway has also patented, for the purpose of raising the heat in the Bessemer converter, a very ingenious modification of the present system. As is well known, a huge flame of carbonic oxide issues from the mouth of the converter, constituting a great loss of heat, as, with but one exception, no attempt has been made to utilise the heat of this flame; but he proposes, by means of a set of upper tuyeres, to burn this carbonic oxide inside the vessel itself. If he succeeds in perfecting such an apparatus, an enormous heat could be maintained, and the use of silicious pigs would cease to be indispensable.

One interesting point deserves notice. Almost invariably, on adding spiegel to the dephosphorised metal, a portion of the phosphorus returns to combination, which must come from the slag, since spiegel contains only minute quantities of that element. Mr. Stead found, some time ago, that spiegel took up phosphorus from slags of this class; that is to say, that manganese reduces free phosphoric acid. To obviate this, it would appear that some modification in the shape of the vessel is desirable to permit of tapping off as much slag as possible.

It is, perhaps, advisable to deal now with the objections which have been raised as to the commercial possibility of this process. They may be briefly summarised as follows:—

1st. Will any and every kind of pig-iron be available? Every possible kind of pig-iron is capable of treatment by this process; but irons very high in silicon are objectionable on account of the extra quantity of basic additions necessary, the increased quantity of slag, and the extra waste and corrosion.

2nd. That whereas the experimental 35 cwt. vessels have answered the purpose, that the operations in the 8-ton vessels may not succeed.

The fact is, that the probabilities of success are just the reverse, since it is far more difficult to retain the fluidity of a small mass than that of a large one, as the sources of heat loss, conduction and radiation, diminish in direct proportion to the mass, or, to use a technical term, the "body of heat" is greater in the larger mass. In actual practice it has been found that the operations in the large converters are, in this respect, quite as satisfactory as those in the experimental vessels.

3rd. That the waste is greatly increased by blowing after the carbon has gone. Mr. Richards stated in London that the waste in the experimental plant was about 17 per cent. Mr. Gilchrist assures the writer that, on a large scale, it is 17 to 18 per cent. As the waste in the ordinary process is about 15 per cent, it is obvious that no very serious difficulty arises from this cause. The amount of slag formed is from 3½ to 5 cwts. per ton of pig-iron used, as against 1½ to 2½ cwts. in the ordinary process.

At the May meeting of the Iron and Steel Institute it was pointed out by the writer, whose opinion was endorsed by Messrs. Stead, Gilchrist, Thomas, and Richards, that "overblowing" (which means oxidation of iron) only commences when all other oxidisable material has been eliminated. The action of the atmospheric oxygen only

commences on the iron when there is nothing else to oxidise. In fact, Bessemer steel-makers seem to forget that the very essence of this process is that there is another element to burn out. Their experience has taught them that when the "flame drops," or, in other words, when the carbon spectrum disappears, that the blow is over, but in the Thomas and Gilchrist process the blow only ends when the phosphorus has gone.

4th. That the corrosive action on the basic lining is greater than that on the ordinary gannister lining, and that while the latter admits of repairs, the former does not.

In reply to this it may be advanced that the basic lining has been repaired, and that Mr. Gilchrist quite recently informed the writer that when using iron not containing excessive silicon that no serious amount of corrosion takes place.

5th. That the result of the process in eliminating phosphorus is uncertain.

In the ordinary process, the disturbing element being absent, there is no danger from this cause, but in this process accidents may occur where the quantity left in the steel may cause difficulty. But in regular working there is very little risk. The following analyses of consecutive blows, kindly supplied by Mr. Gilchrist, show this quite conclusively.

In ten consecutive blows, the samples taken from the converter just before adding spiegel contained—

Phosphorus.									
1.	..	..	..	..	..	..	..	0.10	per cent.
2.	..	..	..	..	..	..	..	0.05	"
3.	..	..	..	..	..	..	..	0.05	"
4.	..	..	..	..	..	..	..	0.07	"
5.	..	..	..	..	..	..	..	0.07	"
6.	..	..	..	..	..	..	..	0.06	"
7.	..	..	..	..	..	..	..	0.14	"
8.	..	..	..	..	..	..	..	0.04	"
9.	..	..	..	..	..	..	..	0.07	"
10.	..	..	..	..	..	..	..	0.10	"

In all cases the pig-iron contained 1.5 per cent of phosphorus. The following are some analyses of steel from the same:—

Phosphorus.	Carbon.
0.11	0.36
0.09	0.21
0.05	0.20
0.08	0.15
0.05	0.08
0.05	0.22
0.04	0.28
0.12	0.40
0.10	0.22
0.07	0.20

Later still, Mr. Gilchrist writes that the last sixteen consecutive blows gave an average of 0.067 per cent P. The C can also be kept at anything from 0.02 to 0.5 per cent, and waste of Mn avoided.

The pieces of steel exhibited were from the Easton Works, and were made by this process. This steel is of excellent quality, though soft. Its analysis is the first of the following; the remaining are analyses of other samples by Messrs. Pattinson and Stead:—

	Rail.	29.	30.	31.	32.
Iron ..	98.944	98.212	98.724	98.743	98.915
Carbon ..	0.350	0.440	0.363	0.385	0.320
*Silicon ..	0.014	nil	nil	nil	nil
Sulphur ..	0.020	0.115	0.093	0.044	0.110
Phosphorus ..	0.062	0.085	0.080	0.058	0.058
Manganese ..	0.610	1.123	0.720	0.750	0.571
Copper ..	—	0.025	0.020	0.020	0.025
	100.000	100.00	100.00	100.00	100.00

* The absence of silicon is a point worthy of note. All analyses of steel made by this process show the same result; for most purposes this fact is decidedly in favour of such steel.

The removal of the phosphorus is not, however, always so complete. M. Pourcel, of Terrenoire, quotes the following analysis of a steel made in his presence:—

	Pig-iron.	Steel.
Carbon	3'20	0'171
Silicon	3'03	traces
Sulphur	0'03	0'037
Phosphorus ..	1'80	0'223
Manganese ..	0'45	0'160

Such a steel as this, although adapted for certain purposes, would not, like the preceding samples, be suitable for best purposes.

6th. The cost of the process.

Sufficient experience has not yet been obtained, but seeing that there is a difference of 10s. or 12s. per ton between the cost of hæmatite and common (*i.e.*, phosphoretic) pig-irons, there is a wide margin to work upon, even in these times of abnormally low prices. The costs may be summed up in the following items:—

- (1.) The slightly increased waste.
- (2.) The extra quantity of slag: the cost of this item mainly consisting in the labour necessary for dealing with it.
- (3.) The cost of the basic additions.
- (4.) The increase in cost of the linings.
- (5.) The decreased productive power of the plant, owing to the facts that smaller charges must be blown, owing to the extra slag formed and risk of splashing, and also the time occupied in taking and examining samples.
- (6.) The smaller production of the blast-furnaces in smelting poor phosphoretic ores as compared with the output of those smelting richer non-phosphoretic hæmatites. But this, the writer contends, is already considered in the price of both pig-irons, and is consequently *nil*.

Taking these objections as a whole, it does not appear that they will amount to anything like 10s. per ton, but possibly 7s. 6d.

As a set-off against these may be considered the utilisation of the large deposits of phosphoretic ores in this and other countries, which may be so much more cheaply worked and delivered to the works than hæmatite ores from distant countries, and the prolongation of the lease of life of inland iron-producing districts in all countries which have their own coal and ironstone. As also the fact, from this latter cause, many existing plants will be as favourably situated as new works built on the coast at a distance from coal-fields, as but few coal-fields of any magnitude are within very easy access of seaports.

There is, however, one objection which may be well left till last, as the weakest of all, and that is, that steel made from such raw materials cannot possibly be so good as that made from pure material. Such an exception has been seriously made.

This objection may be well left to answer itself, since every one must admit that two steels of similar chemical composition, and answering to the same tests, are just as good as one another, even if one is made in a crucible from best Swedish iron and the other in a Bessemer converter from phosphoretic iron. It does not require a spirit of prophecy to justify one in predicting that the day is not far distant when steel made either by the Bessemer or open-hearth process will supersede that made by any other process, and of any quality, however fine it may be.

It may be readily admitted that the Thomas and Gilchrist process has not yet entirely surmounted all its initial difficulties; in fact it would be too sanguine to expect that it could do so in so short a time. It is very probable that no single one of the processes alluded to may be the successful one, but that a combination of several of them may bring about the desired result. Be that as it may, it is obvious that this great problem is solved, and that nothing more is wanting than the rapid and effectual removal of the minor difficulties.

LEAD FUME, WITH A DESCRIPTION OF A NEW PROCESS OF FUME CONDENSING.*

By A. FRENCH.

THIS paper describes a series of experiments made by the author and Messrs. H. J. Wilson and J. Wycliffe Wilson, of the Sheffield Smelting Company, with a view to discover a good process for condensing fumes of lead, silver, and other metals which volatilise in the smelting and refining operations. The loss of lead and frequently of silver by sublimation is an evil with which every smelter is familiar; not only does the loss amount to hundreds of tons of lead in a year at many works, but the injury which is done to health and vegetation is very great. It also describes a new and very successful method of condensing, whereby from 95 to 98 per cent of the metallic contents of the smoke is saved.

The various methods of condensing fumes which have been tried in this and other countries may be classed as follows:—

- (a.) Deposition of the fume by its own gravity in long flues with or without the addition of a series of settling chambers, placed either near to or at some distance from the furnace.
- (b.) Filtering through flues, towers, or chambers containing brushwood, coke, coarsely woven fabric, or similar porous material, using water either in a constant or intermittent stream to keep the filters from becoming choked.
- (c.) The use of water, either in the form of steam, or in showers of drops or jets, projected with some considerable degree of force into and across the current of smoke.
- (d.) Processes based on the inverse of the preceding principle, viz., passing the smoke under and through a depth of water, either in great volumes as in the old Stagg's condenser, or in a more or less comminuted condition.

Our first step was to ascertain the physical nature of the lead fumes and their deportment under varying conditions of temperature and friction. We exposed slips of glass to the fumes for a second or two so as to obtain a very thin deposit on the glass and examined these by a microscope. The first specimen was taken while the lead was in the true state of a gas by holding the glass slip within the flame of burning lead. This when examined by a high magnifying power, presented the appearance of a uniform grey coating, which was perfectly continuous and without any granular structure. A second one taken while dense white smoke was issuing from the tym of the furnace had the same grey film, but in addition it had what appeared to be a superposed layer of small rounded particles or rather spheroids having a considerable degree of oblateness. The sizes of these were very uniform in specimens taken at the same time, although those taken at different times frequently differed greatly. The smallest of them as near as I could measure were about 1-20,000th part of an inch in diameter. A third specimen taken at the mouth of the furnace about ten minutes after charging so as not to catch any particles of dust, had also the continuous film, but much thinner, and the isolated particles were more numerous. A fourth taken from the flue about 60 feet distant from the furnace had no continuous film, but consisted only of the isolated particles, with a few flaky agglomerations of very irregular forms. As the distance from the furnace was increased the flakes became larger and more numerous, but the number of the small particles never seemed to grow less abundant.

It would appear from these experiments that as the vapourised lead cools it assumes the condition of a vast number of minute isolated particles.

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

The state of chemical combination in which the lead in fume is either as sulphate, oxide, or carbonate, generally two or all of these together, and frequently sulphide or sulphite of lead are present in small proportions, and in special operations where chlorides are contained in the furnace charge chloride of lead may be found. Lead fume, especially that from blast-furnaces, nearly always contains a considerable quantity of alumina but very little iron. If there is silver in the ore a portion of it is volatilised with the lead. Although the loss of silver by volatilisation increases with the richness of the furnace charge in that metal, yet the loss does not increase in the same proportion, that is to say, the poorer the ore is in silver the greater is the percentage which is volatilised. This is unfortunate for those who smelt very poor lead ores containing a little silver in blast-furnaces, and produce much fume, for if it were otherwise the smelting of such ores would also be a silver concentration process. For example, a smelting charge containing 30 per cent of lead, while the proportion of silver to lead was only 18½ ozs. per ton yielded fume containing 10½ ozs. of silver per ton of lead. In a second instance, when richer materials were being smelted, the proportion of silver to lead in the charge was 80 ozs. per ton, while the silver in the fume was 33 ozs. per ton of lead. A much richer charge containing 960 ozs. silver to the ton of lead yielded fume containing 100 ozs. silver to the ton of lead. These figures show that as the quantity of silver in the charge increases, the fume increases in richness, but not in the same proportion. I have found the fume at the end of a lead works flue 1000 yards long to contain 5½ ozs. silver per ton, while the proprietor assured me that as far as he knew the ores he smelted never contained more silver than would yield a proportion of 3 ozs. per ton of smelted lead. I only state the facts as far as I know them; the reason I cannot explain.

Lead fume appears to have no definite composition, as the proportions of its constituents vary in every specimen. The lead varies from 35 to 65 per cent. One analysis gave:—

Oxide of lead	44.80
Oxide of zinc	4.80
Oxides of bismuth and copper	1.52
Oxide of iron	trace
Alumina	10.00
Arsenic and antimony (oxides)	3.03
Sulphuric anhydride	28.81
Insoluble silicious matter	9.00
Total	101.96

Another gave:—

Oxide of lead	68.35
Sulphide of lead	2.25
Oxide of zinc	1.80
Lime	2.63
Alumina	5.40
Sulphuric anhydride	16.84
Insoluble silicious residue	2.25
Total	99.52

Lead fume besides silver invariably contains a little gold, usually from ½ to 1 per cent of the quantity of silver.

We have also found on several occasions small quantities of platinum and iridium in the fume, even in that taken from the part of the flues most remote from the furnaces. I cannot believe that the platinum and iridium were carried there in fine dust from the furnace, for in some instances the furnace was smelting rich slags only, which had come from another furnace where matters containing those rare metals were being smelted. I am well aware of the extreme minuteness of the particles of iridium after it has been alloyed with lead and then separated, but in view of the fact that platinum and iridium, especially the latter, have a particularly strong tendency

to unite with sulphur compounds of lead, I incline to the belief that those metals appear in the fume as a true sublimate. We have made many experiments to ascertain what tends most to promote the settling of fume in long flues. We first examined the interior of a flue which extends for several hundred yards in a tortuous course underground. This flue besides having many abrupt turns and angles had partial stoppings of sheet-iron placed at intervals of about 15 or 20 yards. These extended to a height of about 15 inches above the floor, and contracted the area to about four-fifths at those parts. We found that at the various turns and angles, and indeed wherever an eddy of the current occurs the fume lay thickest, in some places heaped up like drifted snow.

The greatest deposition of lead fume takes place as one would expect near the furnace. By following the course of the flue from the furnace to the chimney, we find that the fume lies thickest in the first 100 or 150 yards, and generally beyond that distance it begins to diminish rapidly in quantity, until at about 400 or 500 yards it is only ½ as deep as in the first 100 yards. The relative distances are not alike in every case, but vary slightly according to the kind of furnace employed, the temperature of the smoke and its velocity in the flue. The subsidence of the fume appears to be promoted by whatever causes the isolated particles of which it consists when it leaves the furnace to unite into flaky masses; therefore, the more friction, buffeting, and violent agitation it suffers the more readily it settles. Cooling the smoke through a considerable range of temperature also promotes the subsidence of the fume. This is undoubtedly due to the particles being brought nearer together by the consequent contraction of the gases. I have found that the specific gravity of lead fume from a blast-furnace is about 5.5, and assuming that a cubic foot of smoke contains 4 grs. of fume, which it often does, and that the size of the particles are 1-20,000th part of an inch in diameter, we may find by a simple calculation the number of particles contained in a cubic foot, and their distances apart from each other. Thus:—

1 cubic foot of water weighs 437,500 grs.

1 " " fume " 437,500 × 5.5 = 2,406,250

And as 1 cubic foot of smoke contains 4 grs. fume the aggregate space occupied by the 4 grains will be 4-2,406,250ths part of a cubic foot, or 1-348th part of a cubic inch, but the cubical contents of a sphere 1-20,000th part of an inch diameter is 1-15,278,840,000th of a cubic inch; therefore, dividing 1-348th by that fraction, we get 43,904,712, the number of particles in a cubic foot, and if we extract the cube root of this number it will be the denominator of a fraction, which, having one for its numerator, will express the distances in parts of a foot which the particles are apart, or—

$$3 \sqrt{\frac{1}{43,904,712}} = \frac{1}{352} \text{ part of a foot,}$$

or about 1-30th part of an inch. Hence we learn that those minute particles are about 660 times their own diameter apart, and for the sake of giving a better idea of the comparative remoteness of those fume particles from each other in ordinary lead smoke, if we suppose them to be magnified to the size of the earth, then their distances apart would be 22 times greater than the distance between the earth and the moon.

We invariably find the fume is most abundant wherever the gases have suffered the greatest friction and fall in temperature. That this fact causes the fume to settle is also proved by the increased escape into the air for some time after the flues have been swept out. This I have proved by a great many assays of the smoke, and it is also apparent at the top of the chimney to the eye.

The following experiment proves the extreme difficulty of arresting lead fumes at high temperatures:—We sifted ground slags first through a sieve having 15 meshes to a lineal inch, and the portion which passed through was put upon a second sieve, having 30 meshes to the inch to

remove the dust and finer grains. We then made a filter-bed with this about 1 inch thick, and aspirated a portion of the flue gases through this, taken at two points in the flue, one near the furnaces and another a considerable distance from them. The temperature at the first place was about 750° F., and although the smoke was drawn only at a slow rate the fume passed in considerable quantity, and it continued to escape after a coating of fume about 1-16 of an inch thick had formed at the surface of the filter. In the trial at the distant station, where the temperature was only 300° F., a little fume passed for the first fifteen minutes, and after that the filtration became almost perfect, and the gases then passed only very slowly. This slow passage of the gases through a dry porous filter constitutes the prime difficulty of fume condensing on that principle.

The performance of a long flue is greatly improved by having series of wide settling chambers near, but not quite, at the far end. These prevent the agglutinated particles or flakes from being swept out into the air by the draught. A flue on this principle of about 1200 or 1500 yards, inclusive of about 200 yards of chambers having a width of about two and a half times that of the flue, yields as good results as one 2 or 3 miles long. This is the principle on which the one at the London Lead Company's Works, at Nenthead, in Cumberland, is constructed. The deposition of lead fume may be seen on a small scale very well by aspirating smoke very slowly through a few feet of narrow tube kept cool by water outside, and then through a long wide glass tube placed horizontally and terminating in a wide receiver. A white deposit will be formed along the bottom of the wide tube and receiver, while the upper part, or that which represents the roof of the flue, remains perfectly clean and transparent. We soon learned by experience in our assays of the flue gases that the heavy fume tends to flow along the floor of the flue even when the velocity is as much as 6 or 7 feet per second. This requires to be kept in view when taking samples for examination.

The second class of condensers—namely, filtering through brushwood, cokes, wire netting, tangled wire, or coarse woven fabric—has frequently been attempted, but never with satisfactory results. Such material, if kept dry, does not arrest much fume until they are nearly choked up, and then the draught ceases, and if they are kept wet they scarcely stop any at all.

Our experiments on filtering were chiefly made with wet cokes. In our first we passed the smoke successively through two beds of coke, each 6 inches deep, placed one above the other. A dripping apparatus was attached to keep the cokes wet: the cokes were broken and sifted to the size of from pigeon's eggs to horse beans. The flue gases were propelled through the filter by means of a rotary blower, with a pressure equal to 5 inches of water. This was allowed to run for four days. The smoke which was sent through the filter was drawn off from the main flue at a considerable distance from the furnaces, and contained on an average $\frac{1}{2}$ grain of metallic lead per cubic foot. When the filter was opened the cokes were almost clean; the water in the bottom of the apparatus contained not more than 3 ounces of fume, and after washing the cokes the total quantity did not amount to more than from one-tenth to one-twentieth of the quantity of fume which passed. We afterwards tried the effect of a greater depth of coke. Two towers, 13 feet high and 12 inches wide, were filled with pieces of coke about the size of oranges, and the smoke was sent first up the one and then down the other. An intermittent supply of water was employed to keep the cokes open.

It required a tension of twelve inches water to draw the smoke through the towers. In order to determine accurately the results of our experiments, large samples of the smoke before it entered and after it left the towers were drawn through cylinders filled with cotton-wool, and then through a long glass tube inclined at an angle of about 20°, having a bend at the lower extremity like a cooper's tube, but open at both ends. This contained a little water,

which served to wash the gases after passing through the wool. The gases were drawn through the two sets of apparatus by means of an aspirator worked by steam-jet, and their quantities measured by two large gas meters.

In the above experiments the gases before entering the coke-towers yielded 545 grains metallic lead per 1000 cubic feet, and the gases escaping from the towers gave 356 grains, or 35 per cent caught. A second experiment was made, with a similar result, and as a good deal of power was required to draw the smoke through 26 feet of coke, this method was considered quite useless. It may be well to state that the question of recovering the volatilised silver was of great importance.

We next tried filtering through strong canvas cloth made up into a bag, exposing a surface of about 40 square feet, but this method did not succeed. The fume passed freely through the canvas at first, but after a short time the meshes became completely closed, and the passage of the smoke became so impeded that the experiment had to be stopped. The bag was then washed, and about 3 ozs. of fume were obtained after a run of about an hour and a half.

The following experiment shows with what great freedom lead fume escapes through cloth filters. A glass cylinder, about 14 inches in diameter, was fitted with four calico diaphragms of gradually increasing fineness, the coarser being placed first. After aspirating the smoke through them for an hour they were removed, and each was found to have a perfect coating of fume on the side against the current. Their order was now reversed, the finest being placed first, and using fresh cloth. After running for another hour we found the foremost and finest only had caught fume, and all the others were quite clean.

In other experiments we invariably found that smoke which had passed through a calico filter of a given fineness did not form any deposit on a diaphragm of coarser stuff, but in every case there was an escape of fume through the cloth. This was shown by the darkening of sulphide of ammonium through which the filtered smoke was made to pass.

We now come to the third class of condenser, namely, the use of water either in the form of steam or on the shower-bath principle. One of the earliest attempts to catch lead fumes in this way was made at Allenheads, near Alston Moor. There they had a long flue extending nearly three miles up a hill. Along the first hundred yards two perforated pipes were carried on each side of the flue. Small lead or copper nipples supplied a profuse shower of water. This plan did not succeed well. The small jets got stopped up with bits of straw, grass, and other organic matters, and the quantity of fume caught being no greater than it would have been in a dry flue. The plan was abandoned. The opinion is now very general amongst lead smelters that the drier a flue is the better it arrests the fumes.

The shower-bath principle is now very extensively carried out at the Wanlockhead Lead Works, belonging to the Duke of Buccleugh, in Dumfriesshire. About 2 tons of water are delivered per minute in fine jets over the smoke, while it is ascending and descending in a tortuous course through a series of tall chambers filled with earthenware drain-pipes. Very good results have been obtained from this condenser; that is to say, to obtain large quantities of fume it was found expedient to extend the flue beyond it from time to time until it has now reached a distance of one mile beyond the condenser; and the fact is of considerable importance that about one-half of the fume saved every year is obtained from the long flue, and the other half from the condenser.

Practical results accord well with the theory that rain-drops, with their comparatively enormous surface tensions, are ill adapted to collect the extremely mobile particles of fume they encounter in their descent.

The use of steam in the flues as a condensing agent, although still carried on at some places, scarcely deserves notice as it is positively injurious. Most flues contain an

average of about 5 per cent of moisture, and as soon as this condenses into water there is generally a notable decrease in the quantity of fume which subsides.

The fourth class of condensers consists of those which have for their principle the passing of the smoke through a body of water. This principle has been tried in various ways. The old Stagg's condenser, in which the smoke was drawn in great volumes under the surface of water by means of powerful pumping is now nearly, if not altogether, obsolete.

Our experiments showed that mere bubbling the smoke through water from a perforated pipe, for example, has little effect in stopping the fume. We made experiments to prove this, both on the large and small scale. In the large one we passed the smoke through a number of horizontal perforated pipes submerged 11 inches in water. Our assays showed that only 30 per cent of the fume was arrested. Our experiments on a smaller scale gave even worse results. The reason why simple bubbling through water succeeds no better than the shower-bath principle is that in both cases precisely the same cause operates, viz., the surface-tension of the water, which is just the same whether for a concave or convex surface of equal extent.

We can prove that fume is difficult to wet by coating a glass plate with it, and then dropping water on it while it is held at an angle of about 60° to the horizon: the drops are reflected off without wetting the plate.

This question of surface-tension was well illustrated by an experiment made at the suggestion and in the presence of Mr. Alfred E. Fletcher, one of Her Majesty's Inspectors. Equal quantities of smoke were bubbled through a wash-bottle arrangement, filled first with water, and then with ordinary rape oil. The oil, which has less than half the surface-tension of water, caught more than three times as much fume as the water.

The considerations led us to seek for some way of destroying the surface-tension of the bubbles, and we hit on the device of using fine wire gauze, made of any metal capable of resisting the corrosive action of sulphurous acid. Copper gauze answered perfectly.

In our new apparatus we use wire gauze having about 15 meshes to a lineal inch, the meshes being about 1-20th of an inch wide. A number of gauze diaphragms are arranged one above another in horizontal planes, and at small distances apart. The whole are submerged in water. The smoke is equally distributed under these by means of a horizontal series of perforated pipes. The gauze diaphragms do not add much to the resistance which the smoke current has to overcome in its passage through the apparatus; three diaphragms of the size mentioned above add about $\frac{1}{4}$ an inch of water pressure. The depth of water usually employed is 7 inches above the perforated pipes, and with this depth the water gauge indicates a resistance of about 10 inches, $\frac{1}{4}$ an inch only of which is due to the gauze, the remainder being due to the depth to which the smoke depresses the water at the inlet passages. The ascending gases set up an upward current of water through the gauzes, and to promote a steady circulation of this a return passage is provided.

Although we usually work with three diaphragms of wire gauze, double that number may be used without adding appreciably to the resistance, and by so doing still more perfect results may be obtained. Each square foot of area of the diaphragm space is capable of passing about 40 cubic feet of smoke per minute, and when a blast-furnace is employed for smelting lead ore about 1 foot of area will be required for each ton of ore smelted in twenty-four hours.

During the past six months almost daily assays have been made of the smoke before it entered and after it left the condenser. These have with a few exceptions exceeded 95 per cent of fume caught. The average has been 98 per cent, and in a few cases as high as 99 $\frac{1}{2}$ per cent of the metallic contents of the smoke has been caught. After the lead has been removed from the smoke the large quantity of sulphurous acid which is usually contained in

it may be recovered in a very simple manner. The gas can be mixed with a little air if enough of oxygen is not already present, and then propelled by means of a steam jet through a heating apparatus similar to the hot blast heaters used in iron smelting works, and the hot sulphurous acid steam and air passed through common salt according to Hargreaves's patent process. By this means lead or copper smoke will be rendered not more pernicious than that from ordinary chimneys. Any arsenic or zinc which reaches the condenser is dissolved in the water, and in that way separated from the lead fume, which subsides to the bottom. The apparatus was tried with hydrochloric acid vapour and condensed 97 $\frac{1}{2}$ per cent; of common salt vapour it condensed 93 per cent.

We use a Roots blower, with iron revolvers for forcing the smoke through the apparatus; from 2 $\frac{1}{2}$ to 3 horsepower is amply sufficient to work a condenser large enough for a furnace to smelt 15 tons of lead ore per twenty-four hours. The weight of a condenser for that size of furnace is 18 cwts. The smoke should be cooled to about 120° to 130° F. by passing it through iron pipes, or any other kind of flue. This is necessary to prevent rapid evaporation of the water with which the condenser is supplied. It is very important to cool the smoke as far as possible so as to have a smaller volume to pass, and thereby save both power and cost of a larger apparatus.

THE AMERICAN ASSOCIATION FOR THE ADVANCEMENT OF SCIENCE.

THIS Association held its twenty-eighth meeting at Saratoga Springs, N.Y., beginning August 27th, 1879. The meeting was a memorable one, both on account of the large attendance and the great value of the papers presented. The Association was honoured by the presence of an unusually large number of its Ex-Presidents, no less than nine being on the platform at one time.

The Presiding Officer, Prof. G. F. Barker, M.D., of the University of Pennsylvania, was very happy in conducting the business of the Society, his genial humour in nowise detracting from the dignity of the occasion, and serving as a pleasant refreshment.

The other officers of the Association were as follows:—Vice-President of the Physical Section, Prof. S. P. Langley, of Alleghany, Pa.; Vice-President of the Natural-History Section, Major J. W. Powell, of Washington, D.C.; Permanent Secretary, Prof. F. W. Putnam, of Cambridge, Mass. In the absence of Dr. Geo. Little, of Atlanta, Ga., Dr. H. Carrington Bolton, of Hartford, who was General Secretary at the St. Louis Meeting (1878) was continued in office.

The Chairman of the Subsection of Chemistry, Dr. Ira Remsen, of Baltimore, was unfortunately prevented from attending, and Prof. F. W. Clarke, of Cincinnati, took the position.

One of the features of these meetings is the Address of the Retiring President. On this occasion Prof. O. C. Marsh, of New Haven, had the duty to perform: his subject was "The History and Methods of Palæontological Discovery."

Vice-President Langley gave an Address on "Solar Physics," a subject to which he himself has materially contributed; and Major J. W. Powell gave an Address on "Mythologic Philosophy," having special reference to the mythologies of the Indian tribes.

Dr. Ira Remsen's Address was a plea for the study of Organic Chemistry, a branch which he claims is too often neglected in the courses prescribed in colleges and scientific schools.

These Addresses will appear in full in the annual volume of *Proceedings*.

Heartily welcomed by the meeting was the distinguished astronomer Dr. Otto Struve, Director of the Pulkowa Ob-

servatory, Russia, who is in this country for the purpose of securing a larger object lens for a refracting telescope than has yet been made in the world. The lens is to be made by Alvan Clark and Son, of Cambridge, Mass.

The lion of the gathering was undoubtedly Dr. Thomas A. Edison, of Menlo Park, N.J. He exhibited on Saturday evening, to a delighted audience of 1500 persons, his recently invented Electro-Chemical Telephone. Speech, music, &c., were transmitted from a distant room, and the sounds issuing from the telephone were heard by every individual in the hall.

Perhaps the most remarkable discovery announced was that of Dr. Edison in his paper on "The Phenomena of Heating Metals *in Vacuo* by means of an Electric Current" (*vide* CHEM. NEWS, vol. xl., p. 152), which demonstrated that platinum heated *in vacuo* by electricity becomes denser, harder, more infusible, and less liable to disintegration when heated in a flame. Iron treated in a similar manner becomes as hard as steel and just as elastic. Aluminium melts only at a white heat.

The following is a list of the papers read before the Chemical Section:—

- Action of Ozone upon the Colouring-matters of Plants.—A. R. Leeds.
- Bleaching of Sugar Syrups by Ozone.—A. R. Leeds.
- Reduction of Carbonic Acid by Phosphorus at Ordinary Temperature.—A. R. Leeds.
- Oxidation of Carbonic Oxide by Air over Phosphorus at Ordinary Temperatures.—A. R. Leeds.
- Household Chemistry.—Ellen H. Richards.
- On the Deterioration of Library Bindings.—Wm. Ripley Nichols.
- Observations on the Variations in the Temperature and Chemical Character of the Water of Fresh Pond, Mass.—W. R. Nichols.
- Percentage of Sugar in Sap of the Sugar Maple, and causes which determine its variation; with Note on Pressure of Sap.—Harvey M. Wiley.
- A Modified Method of Collecting and Measuring Gases Soluble in Water.—H. W. Wiley.
- Preliminary Notice of the Revision of the Atomic Weights.—F. W. Clarke.
- On Graphite from the Ducktown Copper Mine.—W. L. Dudley and F. W. Clarke.
- The Great Oberstein Industry: *modus operandi* of Colouring, Cutting, and Polishing Agates and Secondary Gems; illustrated by a fine collection of specimens. Mrs. Erminnie A. Smith.
- Exhibition of Crystals of Sapphire from the Gem Beds of Ceylon, with brief remarks.—A. C. Hamlin.
- Note on a peculiar case of Corrosion of the Metal Tin.—J. W. Osborne.
- Results of Systematic Analyses of Air, designed to discover the cause of Variations in the quantity of Oxygen therein contained.—E. W. Morley.
- On an Accidental Contamination of a Source of Water Supply.—W. R. Nichols.
- On the Limits of Meteorological Conditions governing the extension of Beet-root Culture.—W. McMurtrie.
- A Chemical Examination of Black Ozokerite.—C. Gilbert Wheeler.
- Exhibition of Crystals of Sapphire from the Gem Beds of Ceylon, with brief remarks.—A. C. Hamlin.

The next meeting will be held at Boston on the last Wednesday in August, 1880, under the presidency of the eminent archæologist, Hon. L. H. Morgan, of Rochester.

Sanitary Institute of Great Britain.—The Autumn Congress of this Institute will be held at Croydon, beginning on the 21st of October. There will also be in connection with it an Exhibition of Sanitary Apparatus, Appliances, and Articles of Domestic Use and Economy. Applications for space to be sent to the office of the Curator, Mr. Charles L. Marsh, 138, Fleet Street, London, E.C., not later than Thursday, October 9th.

CORRESPONDENCE.

A NEW METHOD OF PREPARING SULPHURETTED HYDROGEN.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xl., p. 154, Mr. J. Fletcher has described what he calls a new process for the preparation of H_2S by melting sulphur and solid paraffin together. Allow me to state that the process is not new, having been used by myself and others as far back as 1872. I continued to use it for more than twelve months, but found it very unsatisfactory, because of its explosive qualities, and upon that account gave it up. As far as I remember an explosion took place about every seventh experiment; at one time the cork of the flask would be violently ejected and the contents sent to the ceiling, at other times the flask was completely destroyed.

The process works remarkably well except for this one fault, and if Mr. Fletcher has succeeded in overcoming its explosive qualities by the addition of broken tobacco-pipe shanks he has indeed rendered a service to chemists in general.—I am, &c.,

WILLIAM JOHNSTONE, F.I.C., F.C.S.

Wiesbaden, September 29, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Bulletin de la Société Chimique de Paris,
No. 7, 1879.

Observations on a Notice by Prof. Widemann.—M. Schützenberger.—The author rebuts Prof. Widemann's claim to have anticipated his discovery of the existence of an allotropic modification of copper.

Reclamation of Priority on the Subject of a Memoir by M. Stempnewsky on the Preparation of Glycol.—H. Grosheintz.—The author first observed and made known the fact that during the preparation of ethylenic glycol, by the method of Hüfner and Zeller, a certain quantity of bromised ethylen makes its appearance.

Researches on the Incomplete Carbides obtained from American Petroleum.—L. Prunier.—The light petroleum, if submitted to dissociation by heat, become a source of incomplete carbides of different orders, less rich in carbon than the formenic carbide, but capable, if the action of heat is sufficiently prolonged, of re-combining among themselves so as to form more complex compounds. Among the incomplete carbons ethylen, propylen, butylen, and acetylen have been recognised. They are accompanied simultaneously by the ulterior products of their polymerisation, or of their reciprocal combination, benzol, ethyl-acetylen, or crotonyl. By a novel application of the general method of solvents the products have been fractionated according to a principle very different from that of a fractionated distillation and crystallisation. In this manner each of the industrial products may be resolved into a great number of carbides, among which are anthracen, phenanthren, chrysen, pyren, chrysogen, benzerythren, &c. Solvents such as benzol, and even petroleum if applied at a boiling heat and for a long time, fix themselves upon certain of the higher carbides, and bring them down to proportions of carbon not exceeding 94 and 95 per cent.

On Iodised Potassium Iodide.—Antony Guyard.—The author finds that this compound is a true potassium biniodide, though of great instability. Very pure methylic

alcohol, and rendered slightly alkaline, gives with the biniodide of potassium an abundant precipitate of iodoform. Ethylic alcohol under the same circumstances does not give even a trace.

Action of Oxalic Acid upon Chlorates, Bromates, and Iodates.—Antony Guyard.—A boiling supersaturated solution of oxalic acid decomposes the solutions of the chlorates, bromates, and iodates very regularly, with liberation of chlorine, bromine, or iodine respectively, and enables each of these classes of salts to be recognised, not merely because the three elements are easily distinguished, but because the solution takes the colour of the gas or vapour which it contains. The bromates are partially decomposed by oxalic acid, even in the cold, whilst the chlorates and iodates require an elevation of temperature.

Process for Separating and Determining Chlorine, Bromine, and Iodine.—Antony Guyard.—The three bodies should exist in solution in the state of chlorides, bromides, and iodides. If wholly or partly present as chlorates, &c., they must be reduced by treatment with an excess of sulphurous acid. The mixture, acidulated with sulphurous acid, is treated with a slight excess of a mixture of bisulphite of soda and sulphate of copper. The iodine is precipitated immediately and very completely as cuprous iodide, in which state it may be directly determined with much accuracy. The results are liable to be erroneous only in presence of sulpho-cyanides—a circumstance not to be expected in practice. After filtering off the cuprous iodide the liquid is boiled with an excess of sulphuric acid until all the sulphurous acid of the sulphites has been completely expelled. When this is effected the liquid is introduced into a flask which communicates, by means of a bent tube, with a Will and Varrentrapp's nitrogen-tube. Into this is introduced a solution of bisulphite, or of sulphurous acid, or bisulphide of carbon, or, if it is preferred, a solution of potassium iodide, and the tube is kept in cold water. To the liquid in the tube is added a small excess of pure chromic acid, or a mixture of sulphuric acid and potassic bichromate, and the liquid is boiled until the bromine is completely expelled. It is then determined, either as bromide of silver, or by a colorimetric process, or indirectly by the volumetric determination of the iodine which it liberates. To determine the chlorine it is merely needful to reduce the excess of chromic acid by means of a sulphite, and to precipitate with silver nitrite.

Decomposition of Hydracids by Metals.—M. Berthelot.

Reciprocal Displacements between Oxygen, Sulphur, and the Halogens combined with Hydrogen.—M. Berthelot.—These two papers have been already noticed.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 67, July, 1879.

Report by M. de Luynes on M. C. Lorilleux's Manufacture of Printing and Lithographic Inks.—Two kinds of varnish are employed in the manufacture of printing inks; the one obtained by boiling linseed oils, and the other from a mixture of resin and resin oil, the latter being chiefly used for newspapers where rapid drying is of importance. M. Lorilleux allows his linseed oil to rest for two years at a constant temperature. It is then boiled by means of hot air, at a distance from the furnaces so as to remove every risk of fire. A mass of 2500 kilos. is boiled from twenty-four to fifty-six hours, and is stirred by a mechanical agitator. The varnish thus obtained is limpid and flows well. The lamp-black is produced either by means of specially constructed lamps or by the decomposition of naphthalin oils, which fall by drops into a heated retort. The gaseous products are carried off by tubes, at the end of which they are burnt

under sheet-iron bells, whilst the black is carried off by a current of air into large chambers. It is afterwards submitted to calcination. All the inks, lithographic or typographic, are submitted to a practical trial before being sent out.

Biedermann's Central-blatt für Agrikultur-Chemie.
August, 1879.

Researches on Damage to the Soil and the Crops by the Waste Waters and the Gases from Manufactures.—Dr. J. König.—These researches relate, in the first place, to the waters from certain mines of zinc-blende in Westphalia, which pass into certain streams used for irrigation, and seriously injure the productiveness of the soil. The presence of zinc oxide in the earth is indicated by the prevalence of *Viola calaminaria*, which contains in its ash as much as 21 per cent zinc oxide. The author has likewise examined the waste waters from a dye works, a wire works, and from pyrites washing. The two latter he considers as directly poisonous to plants on account of their percentage of ferrous sulphate. The dye water may, by reason of the organic matters which it holds in solution or suspension, gradually overload the soil with humus and render it boggy.

The Transformations of Silicates.—Dr. J. Lemberg.—In the first section of this dissertation the author gives a series of analyses of minerals in various stages of transformation and weathering. In the second, he takes up the questions first raised by Berthollet concerning the influence of masses in chemical action. His experiments confirm the views of the French chemist. In the reactions of common salt upon gypsum, of alkaline and magnesian chlorides upon calcium carbonate, there are formed not merely two new compounds by double decomposition but all the four compounds possible in such cases are produced in quantities which vary with the proportions of the salts originally employed. In the third section the author contends that basic feldspars weather easily, but those of an acid character with difficulty. In his fourth section he examines whether in the absorptive action of soils peculiar forces come into play, as Liebig maintains, or whether, as Way and Mulder suppose, the phenomena are due to purely chemical action. In the fifth and last section the author describes experiments in which silicates were added to melted chlorides of potassium, sodium, and calcium, and to sodium sulphate, and gives an account of the changes observed.

Researches on the "Clover Sickness" of Soils.—Dr. A. Emmerling and Dr. R. Wagner.—A soil of this kind was found deficient in phosphoric acid and especially in potash.

Experiments on the Excretion of Gaseous Nitrogen from the Albuminates Transformed in the Animal System.—J. Seegen and J. Nowak.—The authors conclude from their experiments that the animal organism is capable of excreting in the gaseous state a part of the nitrogen liberated on the transformation of the albuminates.

Reimann's Färber Zeitung,
No. 31, 1879.

This issue contains nothing of general interest.

No. 32, 1879.

Vegetable Silk.—A. Müller.—The author, after referring to the fact established by Mulder, that silk when dissolved in cupric oxide, glycerin, &c., is not decomposed, and can be precipitated unchanged, chemically speaking, mentions a patent which he obtained in 1871 for dissolving silk waste and precipitating it upon vegetable fibre.

No. 33, 1879.

A great part of this issue is taken up with complaints concerning the unremunerative state of the dyeing business in various parts of Germany.

A serious, probably fatal, explosion of benzin (benzol or light petroleum-spirit?) took place in a dye-house at Blankenburg. The cause of the calamity is not specified.

No. 34, 1879.

The spinning works of Steinbrecher in Trübau have been since last year lighted up with the electric light. The dye-houses in Spindler's establishment at Spindlersfeld are about to be illuminated in the same manner.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 8, 1879.

Determination of Atomic Weights and the Utilization of Isomorphism for this Purpose.—Hermann Kopp.—This important memoir, which extends to upwards of fifty pages, is unfortunately not capable of useful abstraction.

New Determination of Manganese with the Application of Volhard's Process for the Titration of Silver.—C. Rössler.—The solution containing the manganese in the state of a proto-salt, free from chlorine or other halogen and from organic matter, is placed in a flask holding a half or a quarter litre. A measured quantity of a solution of silver of known strength (10.8 grms. silver per litre) is then added, the quantity being greater than what is requisite for the precipitation of the manganese. The flask is then heated in the water-bath, and a solution of carbonate of soda is added, till not only the black compound but the excess of silver is perfectly precipitated. Ammonia is then added, avoiding excess (about 10 c.c., at sp. gr. 0.958, to every 50 c.c. of the silver solution); the flask is shaken, cooled, filled with water up to the mark, and well shaken. The contents are then filtered through a folded filter into a dry beaker, and the silver present in a measured volume is determined by Volhard's sulphocyanide process, the liquid having been previously acidulated with nitric acid. The quantity of silver solution needed for the precipitation of the manganese present is then found by calculation, 1 c.c. representing 0.00275 gm. of manganese.

Percentage of Sulphuric Acid in Wine.—G. Lunge.—The author points out that Prof. Nessler, led astray by the misprint "centigrammes" for "milligrammes," in an essay by Marty in the *Journal de Pharm. et Chimie*, has declared that no wine ought to be condemned as plastered unless it contains per litre more than 3.28 grms. $\text{SO}_4\text{H}_2 = 5.83$ grms. potassium sulphate. The serious nature of the error will appear on remembering that Prof. Nessler himself only once met with a sample of natural wine containing as much as 0.063 per cent $\text{SO}_3 = 1.36$ grms. potassium sulphate per litre.

New Colouring-matters.—Otto N. Witt.—If the solutions of molecular quantities of meta-toluylen-diamin and nitroso-dimethyl-anilin hydrophlorate in warm water are mixed, there is formed a deep blue liquid, from which a colouring-matter can be precipitated by means of salt. The substance obtained, $\text{C}_{15}\text{H}_{18}\text{N}_4\text{HCl} + \text{H}_2\text{O}$, dissolves readily with a bright blue colour in cold water, alcohol, and glacial acetic acid. By the use, on the one hand, of ortho- and para-toluydin, of the isomeric toluylin-diamin and xylinin, and, on the other, of nitroso-dimethyl-anilin, nitroso-phenol, and of the other known nitroso-compounds, the author has obtained a number of other new colours, with whose investigations he is now engaged.

On Phosphorus Sulphides.—G. Ramme.—The author obtained the penta-sulphide from a solution of sulphur and phosphorus in carbon sulphide. The preparation of the trisulphide in the same manner did not succeed, nor was P_4S_3 (Lemoine) formed.

Artificial Atropin.—A. Ladenburg.—The author has effected the synthesis of atropin from its decomposition products, tropin and tropic acid.

On Tropidin.—A. Ladenburg.—In his investigations of tropin the author has discovered a new base, tropidin, which appears to have a considerable analogy with coniin. He thinks it possible to obtain coniin from tropidin by the addition of hydrogen.

Certain Derivatives of Tropic Acid.—A. Ladenburg.—The derivatives in question are tropide, tropic ether, and chlorhydro-tropic acid.

On Di-isobutylamin.—A. Ladenburg.—The author's observations differ from those of Reimer.

Derivatives of Ortho-toluylen-diamin.—A. Ladenburg and L. Rügheimer.—The authors have studied the behaviour of ortho-toluyden-diamin with acetophenon and with acetic ether.

On Cuprous Chloride.—M. Rosenfeld.—It is commonly stated that this compound is converted into copper sulphate by the action of concentrated nitric acid. In the cold, however, there is no action, and even with the aid of heat very little. Partially oxidised cuprous chloride can be rendered colourless by washing in glacial acetic acid. In dilute nitric acid cuprous chloride remains colourless if light be excluded. It is exceedingly susceptible to light. The author describes certain chromates of copper, with whose further investigation he is engaged.

Action of Chlorine upon Naphthalin- β -sulphon-chloride and a New Trichlor-naphthalin.—Oscar Widmann.—A description of the tetra-chloride of naphthalin- β -sulphon-chloride, of dichlor-naphthalin- β -sulphon-chloride, and of β -trichlor-naphthalin.

Dichlor-naphthalin- β -sulphonic Acid and its Salts.—O. Widmann.—The acid in question is powerful and expels carbonic acid from its salts. Its potassium, ammonium, silver, barium, lead, manganese, zinc, and copper salts are described.

On Nitroso-sulph-hydantoin.—R. Maly.—Nitroso-sulph-hydantoin is characterised by forming with bases bright yellow, orange, and red compounds, which are not true salts.

Decomposition of Ammonium Formiate at High Temperatures.—R. Andreasch.—The chief decomposition-product of formiate of ammonia is not hydrocyanic acid, but formamide. Of the former body only traces are perceptible.

A New Base, $\text{C}_{10}\text{H}_{18}\text{N}_2$.—C. Böttinger.—This base is obtained as a product of the reaction of benzol-chloride with aniline. On heating the base with mercuric chloride or with arsenic acid it is converted into a true colouring-matter. In the former case the solution is first clear, and then reddens, the colouring-matter being developed with a brisk reaction. The crude dye is a reddish violet.

Action of Ammonia upon Quinons.—E. v. Sommaruga.—The author has made a number of experiments on the action of ammonia upon the reduction-products of the quinons. From dioxy-indol, the reduction-product of isatin, he has isolated two colourless crystalline bodies and a fine crystalline red dye.

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THE CHEMICAL NEWS.

VOL. XL.



ON THE
NON-EXISTENCE OF NASCENT HYDROGEN.

REDUCTION OF POTASSIUM PERCHLORATE.

By Dr. D. TOMMASI.

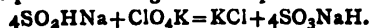
A SOLUTION of chemically pure potassium perchlorate was submitted to the action of various reducing agents, and the following results were obtained:—

(1.) The perchlorate treated with zinc and dilute sulphuric acid at the ordinary temperature, is not transformed into chloride, even by the aid of heat.

(2.) Magnesium and dilute sulphuric acid do not reduce the perchlorate, either in the cold or by the aid of heat.

(3.) A certain quantity of perchlorate was dissolved in a concentrated and boiling solution of copper sulphate. Into this solution a piece of zinc was plunged, and immediately a violent reaction took place, and hydrogen was freely evolved, but the perchlorate underwent no reduction. Neither does any reduction take place when the perchlorate is treated with sodium amalgam, zinc and potash by the aid of heat, &c., nor, indeed, with any of the principal reducing agents in ordinary use.

Now, this perchlorate which undergoes no reduction, although found in the presence of *nascent hydrogen*, is easily transformed into chloride by the action of a compound which does not set hydrogen free, viz., sodium hydrosulphite. The action of sodium hydrosulphite doubtless takes place according to this equation:—



During the action of zinc on sodium bisulphite no hydrogen is evolved, but let us, on the contrary, admit that some is produced. How, then, can it be explained that this same perchlorate, which undergoes no reduction by means of nascent hydrogen, as shown by sixteen different reactions, can be reduced by the hydrogen disengaged by the action of zinc on sodium bisulphite.

M. Würtz, in his *Atomic Theory*, declares himself to be in favour of the nascent state of bodies, and he explains it in these terms:—

"The special activity of hydrogen and oxygen gas in the nascent state is without doubt due to the circumstance that in this case the atoms act separately before being united to a congener in order to form the couples of which the molecules are composed. It is evident that this formation, which is a combination, ought to evolve heat. The separate atoms which are nascent, and not yet united to their fellows, are still provided with that heat, and are, as a consequence, so much the more active." That the atoms of hydrogen are more active than its molecule we do not deny. It appears to us, however, unlikely that when the hydrogen is set free by a reaction, it can be in the state of *isolated atoms*.

We know that copper, even finely divided, is very slightly attacked by hydrochloric acid at the ordinary temperature. Copper hydride, on the contrary, is decomposed very energetically.

"How can this fact be explained," justly remarks M. Würtz, "if, to the affinity of chlorine for copper, be not added the affinity of the two atoms of hydrogen to form a molecule?"

But how are we then to understand that the hydrogen set free by this reaction can be found in the state of isolated atoms, since it is the very affinity of the two atoms of hydrogen that determines the reaction between the hydrochloric acid and the copper hydride?

What has just been said for copper hydride and hydrochloric acid may be said equally for all the reactions producing hydrogen,—say, for example, the action of hydrochloric acid on zinc—



In this case, as we know, there neither is nor can be any hydrogen in the state of *isolated atoms*, as M. Würtz thinks, and the special properties of nascent hydrogen can be attributed only to the calories which accompany hydrogen while it is being set free.

Are we therefore to conclude that hydrogen can be active only in the molecular state? Certainly not; for hundreds of examples prove to us, on the contrary, that in many cases it is not the molecule of hydrogen that acts, but its atom.

An isolated atom cannot exist, and consequently it is devoid of properties. For an atom to exist it would be necessary for its molecule to contain only a single atom, as in the case of mercury, which in all probability is confounded with its atom.

Dr. J. H. Gladstone and Mr. A. Tribe have recently found that there is a great analogy between the effects produced by nascent hydrogen, occluded hydrogen, and those obtained by the copper-zinc couple.* We are happy to find that the result of their researches coincides entirely with our hypothesis, which, as is well known, considers the different allotropic states of hydrogen as ordinary hydrogen in different physical conditions.†

Laboratoire de Chimie de l'Institut des
Hautes études, Florence.

SOME EXPERIMENTS WITH VARTRY WATER.

By JOSEPH FLETCHER, F.C.S.,
Membre de la Société Chimique de Paris, &c.

THERE is a Royal Commission now sitting to investigate the causes of the excessive death rate in Dublin, and although its principal business is with the public drainage and house sanitation, still the results of recent experiments with the water supply of the city may not be unwelcome to some of your readers who take an interest in such matters.

The City of Dublin, and several of the outlying townships, are supplied from an artificial reservoir formed by damming the valley of the River Vartry. The works are on a large scale, and the supply is very abundant, allowing of a constant service at high pressure, besides a very large margin for waste. It is a water which upon analysis shows very little chlorine, 0.001155 per litre, or 0.8025 grains per gallon, which quantity is remarkably constant. It is of great softness, the hardness being only 3 degrees on Clark's scale, and yielding a total solid residue, varying, as the result of many experiments, from 4 grains to 6 grains per gallon.

Two questions have been raised about this water. One is that it is largely contaminated with peat, and the other that its action upon lead is considerable. My experiments were made to throw some light upon these points.

The Vartry is admittedly a peaty water, purified more or less by filtration; but in the early mornings, when the service taps are first opened, it is often of a coffee colour, and vegetable filaments are abundant. I rejected such, and took my samples at mid-day, when the water was uniformly clear. I have found an excellent means of observing the colour is to employ an inverted glass fern shade fixed upon a suitable stand, and filled with the water to be examined. It requires a good many litres to fill it, but where waters are studied at their source that does not matter. Samples which in the quart decanter and the 2-foot tube appear clear, or at most faintly turbid, when

* *Journal of the Chemical Society*, April, 1879; *CHEMICAL NEWS*, vol. xxxix., p. 91.

† *Le Monde*, September 11, 1879.

viewed at a proper angle (from above through the concavity at the side of the apparatus) show a decidedly peaty tinge. This is very evident in the Vartry water. 500 c.c. of the water evaporated gently to 50 c.c. presents the appearance of a strong solution of peat, large flocculi floating in the fluid: filtered, they remain upon the filter, and the filtrate showed a corresponding reduction of solid residue.

My principal object, however, was to investigate the action of the water upon lead. The pieces of metal employed were rolled so as to fit easily into a glass tube connected with the main supply. They were carefully washed in distilled water, dried in an air-bath at low temperature before and after the experiments, and handled as little as possible. The pressure at the tap was unfortunately not regular, which will account for some apparent discrepancies, but this will be arranged in future experiments.

A piece of lead 12.7×7.6 centimetres, placed in the tube for 36 hours lost 70 milligrams; in another 12 hours it again lost 70 milligrams; 48 hours, nothing; 30 hours, 40 milligrams; total loss in 126 hours, 180 milligrams.

A second piece, 8.5×5 centimetres—After 26 hours lost 15 milligrams; in another 22 hours it further lost 40 milligrams; 48 hours, 15 milligrams; 36 hours, 40 milligrams; total loss in 132 hours, 110 milligrams.

The piece employed in experiments No. 2 was cleaned and polished—After 32 hours it lost 5 milligrams; in another 26 hours it lost 10 milligrams; 48 hours, 10 milligrams; 60 hours, 20 milligrams; total loss in 166 hours, 45 milligrams.

Several experiments were then made with pieces of lead in still water, the principal being 14×2.5 centimetres.

After 24 hours it gained 10 milligrams; after another 27 hours it gained another 10 milligrams; and after another 56 hours it remained without change.

The same piece, taken from the still water and placed in the tube—After 48 hours it lost 20 milligrams; after another 24 hours it lost another 10 milligrams; total loss in 72 hours, 30 milligrams.

Again, a piece of lead 7.5 centimetres square was placed in a glass dish, into one end of which a stream of water was allowed to flow, but not in immediate contact with the current: there was no change after four days.

The results of these experiments indicate a water of great purity, chemically considered, but strongly impregnated with peat, having a very decided action upon lead when flowing through pipes of that material, but without action upon it when at rest, but rather leaving an organic deposit.

I am making further experiments upon this water, which I shall have pleasure in submitting.

ON SOME POINTS IN CONNECTION WITH AGRICULTURAL CHEMISTRY.

At the recent meeting of the British Association for the Advancement of Science, held at Sheffield, Dr. Gilbert, at the request of the President of the Chemical Section, gave some account of results obtained at Rothamsted, of which the following is briefly the substance:—He stated that, in the course of the experiments conducted there, wheat had now been grown for thirty-six years in succession on the same land, barley for twenty-eight years, oats for nine years, root-crops for more than thirty years, and beans also for about thirty years. Experiments on an actual course of rotation had extended over thirty-two years; and, lastly, experiments had been conducted on the mixed herbage of grass land, in Mr. Lawes's Park, for twenty-four years. They found minor distinctions in the manurial

requirements of different plants of the same natural family, but very great distinctions in the requirements of plants of different natural families. The gramineous crops are very low in their percentage of nitrogen, and yield but a small quantity of it per acre. Yet nitrogenous manures are very effective when applied to such crops. Leguminous crops, on the other hand, are very high in the percentage of nitrogen, and yield a large amount of it per acre. Yet nitrogenous manures are of little avail to those plants, and potash manure is especially effective. The difference in the manure-requirements of plants of some other natural families was also pointed out.

Much more complicated, however, was the problem, when experiments were made upon the mixed herbage of grass land, where they might have fifty or more species growing in association, representing perhaps twenty natural families. It was at once found that the manures which most favoured gramineous crops, grown separately on arable land, brought forward the gramineous plants in the mixed herbage. Those, on the other hand, which favoured the Leguminosæ, grown separately on arable land, brought forward the Leguminosæ in the mixed herbage. The plants of other natural families also exhibited characteristic susceptibility to the manures employed. In fact, any manure—that is, anything that increases the growth of any species—induces a struggle, greater or less in degree, causing a greater or less diminution, or a disappearance, of some other species. Hence the twenty different plots, in the experiments in question, soon showed as many distinct floræ. Tables were exhibited illustrating the variation in the number of species, which was in some cases fifty and in others under twenty; and the percentage by weight, and the amounts per acre, which the different natural families yielded, were also shown.

There were very great differences, not only in the flora, but also in the character of development of the plants, degree of luxuriance, tendency to form leaf or stem, to mature or otherwise, and so on; and, with these, there were also very great differences in the chemical composition of the produce. The dry matter of the mixed herbage contained, per cent, in some cases $1\frac{1}{2}$ times as much nitrogen as in others; and the amount of nitrogen in the produce per acre was three times as much in some cases as in others. The percentage of potash in the produce varied as one to two, and the amount of potash yielded per acre as one to five, in the different experiments, and there were considerable differences among the other constituents. The produce of the respective natural families, when normally developed and when ripe, may be said to possess a characteristic composition within certain limits. Yet the composition varied immensely according to the conditions supplied, the species grown, and the character of development induced. Thus, the ash of the separated gramineous produce showed a variation in the percentage of potash of from about 24 to about 40; the ash of the leguminous produce, from 12 to 33; and that of the mixed produce of the other natural families, from 17 to 37.

One point of especial interest was the difference in the amount of nitrogen taken up over a given area by plants of different natural families. The measurable, or as yet measured, annual deposition of combined nitrogen from the atmosphere was quite inadequate to account for the amounts taken up by the vegetation. Even unmanured land may lose more than this by drainage. It was assumed by some that some plants assimilated the free nitrogen of the atmosphere, whilst others did not; and if this were established many existing difficulties would be explained away. But Mr. Lawes and he (Dr. Gilbert) considered that the balance of the direct experimental evidence on the point was decidedly against the supposition of the assimilation of free nitrogen. The balance of existing indirect evidence was also in favour of the supposition that the different plants only took up combined nitrogen, and chiefly from the soil. It was shown, by reference to their experiments at Rothamsted, that in the growth of wheat or barley for many years in succession

on the same land, without nitrogenous manure, the annual yield of nitrogen in the crop gradually diminished. With this there was a diminution in the percentage of nitrogen in the soil. In the case of the root-crops, where, under similar conditions, the diminution in the annual yield of nitrogen was greater than in the case of the cereals, the diminution in the percentage of the nitrogen in the soil was also greater. In the case of beans there was also a diminution in the yield of nitrogen in the crop, but still much more was yielded over the later period than in either wheat or barley. The bean-field did not, however, show a marked reduction of nitrogen in the surface soil. In the case of the mixed herbage experiments very much more nitrogen was yielded by the application of potash manure, in great part due to the increased growth of leguminous plants; and here they found a great reduction in the percentage of nitrogen in the soil. In the case of clover, grown for many years in garden soil, the percentage of nitrogen in the soil was also very largely reduced. Part of this reduction might be due to loss by drainage, and in other ways; but the indication was that the Leguminosæ had derived their nitrogen from the soil.

Admitting that the sources of the whole of the nitrogen of vegetation were not conclusively made out, they nevertheless considered that the existing evidence was against the idea of the assimilation of free nitrogen by plants, and in favour of the opinion that the nitrogen was mainly, if not entirely, derived through the medium of the soil. Independently of the combined nitrogen which soils receive from the atmosphere, or by manure, most have large accumulated stores derived from past ages of vegetation, with perhaps greater normal annual supplies than at present, and certainly less removal, and therefore gradual accumulation. On this point it may be mentioned that a sample of Oxford clay, obtained in the recent Sub-Wealden exploration boring, at a depth of between 500 and 600 ft., showed, on analysis at Rothamsted, approximately the same percentage of nitrogen as the subsoil at Rothamsted taken to a depth of about 4 feet only. An acre of moderately clayey soil and subsoil may indeed contain several thousand pounds of combined nitrogen within the depth to which the roots of growing crops descend.

ON THE FORMATION OF ROSANILINE.

By JUSTUS WOLFF.

THE chemical process of the formation of rosaniline is much more complicated than it appears, and it is worth while to examine it in all its details.

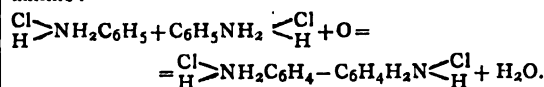
In order to manufacture magenta, a suitable mixture of aniline and toluidine is treated at an elevated temperature with an oxidising or dehydrogenating agent, which is an acid by itself, or with a neutral or basic oxidiser or dehydrogenator in presence of an acid. The presence of an acid is one of the first conditions for the formation of rosaniline: without acid no rosaniline can be formed. The action of the oxidiser or dehydrogenator takes place at a temperature equal to the boiling-points of aniline or toluidine, or their mixtures.

Aniline possessing the lowest boiling-point in that mixture will be acted upon in the first instance, until the boiling-point of a mixture of 2 molecules of aniline and 1 molecule of toluidine is reached, whereafter that mixture will be dehydrogenated, and so on. In the aniline salt, which is acted upon by oxygen or any other dehydrogenator, the nitrogen is in a pentadic state, and therefore its nucleus forms with the acid a stable compound, which is not dissociated at the temperature of the boiling-point of aniline, consequently the oxidiser can act only on the phenyl nucleus of the aniline, eliminating one hydrogen.

Oxygen, being a dyad, requires two hydrogens, and therefore one oxygen will act on the phenyl-nuclei of two molecules of aniline salt, eliminating one hydrogen of

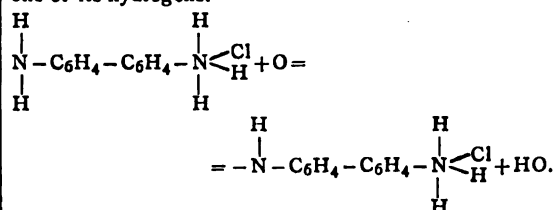
each phenyl nucleus, and producing in each of them one free atomicity, by which they combine together.

For the sake of simplicity I take in the following formulæ hydrochloric acid as the acid combined with aniline:—

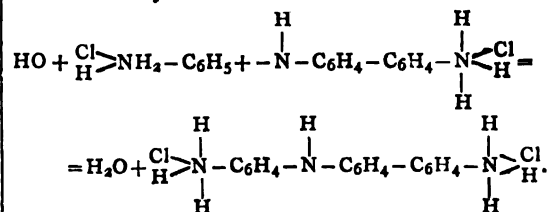


The resulting molecule represents an aniline, in which one hydrogen is substituted by the complex $\text{C}_6\text{H}_4\text{NH}_2$, and therefore it can saturate only one acid molecule, especially at the high temperature of 182° .

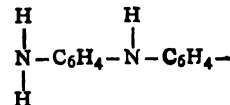
On that account the acid contained in the dehydrogenated aniline, which is substituting one hydrogen in the original aniline, will separate from its nitrogen appendix, whereby the latter is exposed to the action of the oxidiser and loses one of its hydrogens.



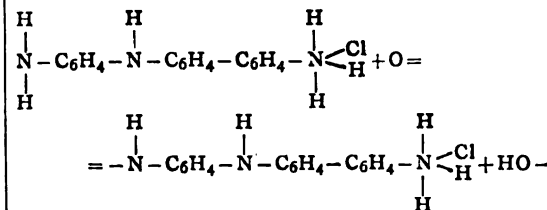
Here we have a compound molecule with one free affinity, besides an oxygen compound with a free affinity too. The latter acts on a new molecule of aniline salt, eliminating one hydrogen of its nucleus, and thus produces a compound with a free affinity, by which it will combine directly with the above molecule.



The last-formed molecule represents an aniline, in which one phenyl nucleus hydrogen is substituted by the complex—

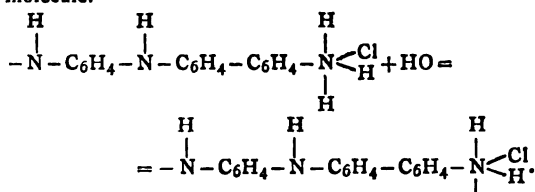


As such it is not able to combine with more than one acid molecule, especially at the high temperature of 182° , and therefore it will separate with the last added acid molecule, by which reaction a change of atomicity of the next situated nitrogen is produced, and consequently the respective nitrogen appendix dehydrogenated—

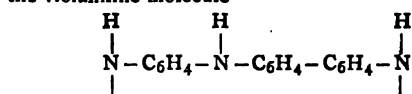


In this complex the acid is completely saturated, and is not able to take up another phenylene compound, and therefore the hydroxyl will not act upon a fresh molecule of aniline salt but on the above compound, and will eliminate one hydrogen of it, which latter is offered for this

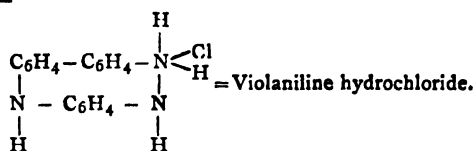
action in the nitrogen appendix NH_2 , joined to the acid molecule.



The last given compound has two free atomicities in its nitrogen appendices, which therefore will combine together, and thus form a closed chain or ring, representing the violaniline molecule—

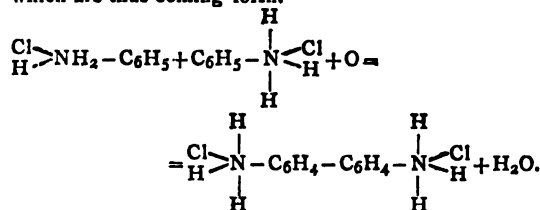


Or—

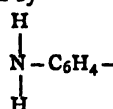


The formation of violaniline starts at a temperature equal to the boiling-point of aniline, and ends as soon as the boiling-point of a mixture of two molecules aniline and one molecule toluidine is reached, viz., 188° . From 188° upwards begins the formation of mauvaniline in a manner similar to that of violaniline.

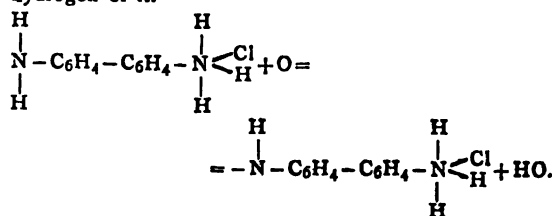
In such a mixture the oxygen acts in the first instance on that part which possesses the lowest boiling-point, consequently on the aniline. Two molecules of aniline being present, the oxygen acts on the two phenyl nuclei of them by eliminating one hydrogen of each, in consequence of which they combine together by their free atomicities, which are thus coming forth.



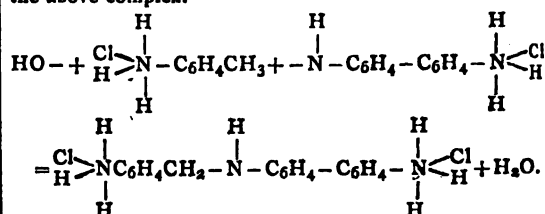
This compound, representing an aniline, in which one hydrogen is substituted by—



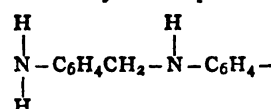
can therefore combine with only one acid, and on that account has to separate with the other acid molecule. In that manner a change of atomicity takes place in the nitrogen, and in consequence of it this nitrogen appendix will be acted upon by the dehydrogenator, eliminating one hydrogen of it.



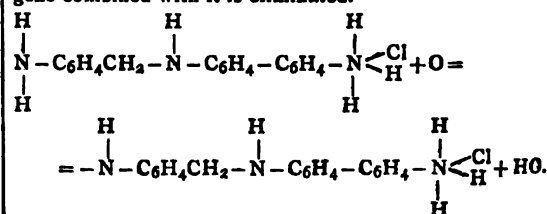
The last compound, not being able to saturate the acid molecule completely, and possessing one free atomicity in presence of a dehydrogenator, the latter will act on a fresh molecule, which in this case is toluidine, and eliminate one hydrogen of its methylene appendix, thus creating a free atomicity in it, by which the toluene compound joins the above complex.



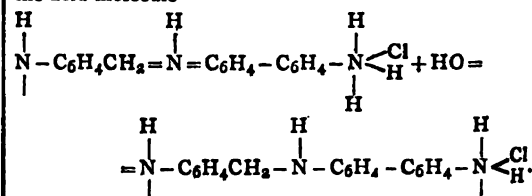
This last compound represents an aniline, in which one hydrogen is substituted by the complex—



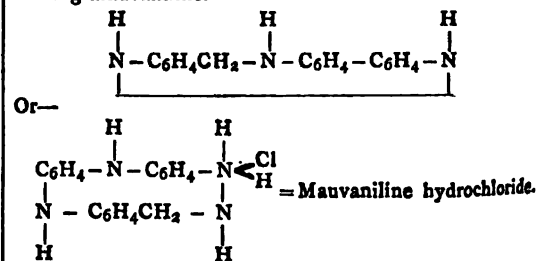
and which therefore can combine with only one acid molecule, and on that account it has to separate with its second acid molecule. By this action the nitrogen next to this second acid molecule changes its atomicity, and is subsequently acted upon by the oxygen, and one of the hydrogens combined with it is eliminated.



In that last compound the acid molecule is completely saturated, and therefore the compound cannot take up another phenylene nucleus, on account of which the hydroxyl does not act on a fresh aniline or toluidine salt, but on the compound itself, by eliminating a hydrogen, which can only be that in the NH_2 appendix, nearest to the acid molecule—



Both free atomicities of the extreme nitrogens join themselves together, thus forming a closed chain, representing mauvaniline.

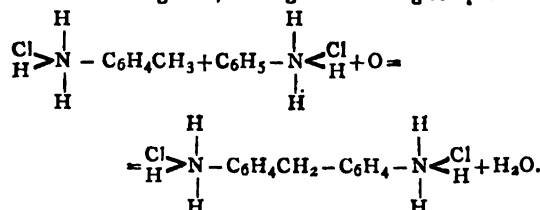


As stated above, the formation of mauvaniline begins at about 188° , and goes on until the boiling-point of a mix-

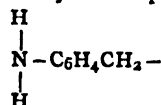
ture of one molecule of aniline and two molecules of toluidine, viz., 193°, is reached, at which point commences the formation of rosaniline.

In treating a mixture of one molecule of aniline and two molecules of toluidine with an oxidising or dehydrogenating agent, the lower boiling constituent will be acted upon first, and this is in that case the aniline which by that action loses one of its phenyl nucleus hydrogens, whilst the oxygen is converted into hydroxyl, acting on a fresh molecule in the mixture; and in this case toluidine eliminating one hydrogen of the latter's methylene appendix.

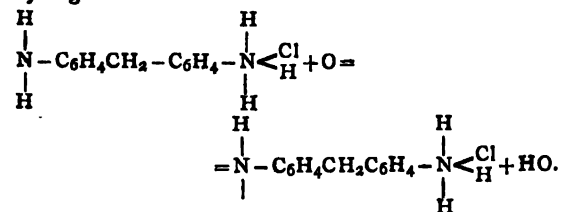
Thus we have a phenylene-amine with a free atomicity, and a toluene-amine with a free atomicity, by which they both combine together, forming the following compound:—



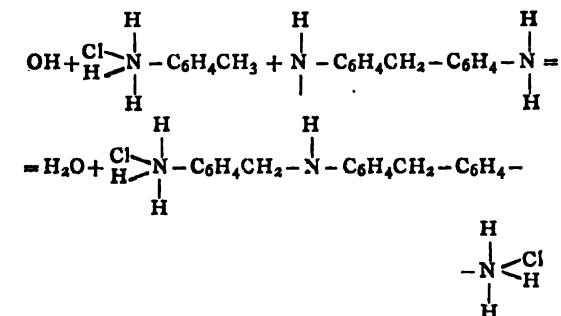
This compound representing an aniline, in which one hydrogen is substituted by the complex—



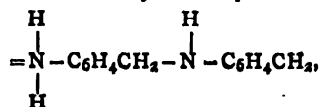
can combine only with one acid molecule, and therefore separates with the second one, leaving its nitrogen appendix to the action of oxygen, which latter eliminates one hydrogen.



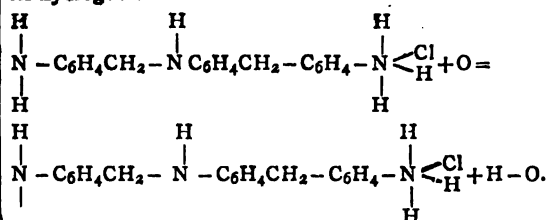
The hydroxyl coming forth by this process acts on a fresh molecule of toluidine, eliminating one of its methylene hydrogens, by which a free atomicity is produced. By that free atomicity the formed toluene amine hydrochloride combines with the last derivated molecule:—



The latter complex representing aniline in which one hydrogen is substituted by the complex—



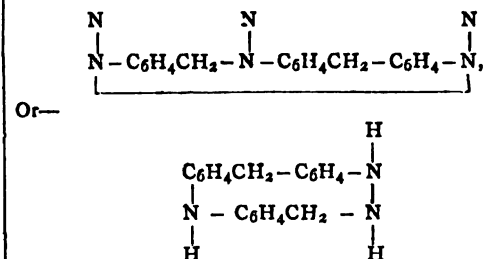
which can combine with only one acid molecule, will separate with its second acid molecule, and thus expose its nitrogen appendix, nearest to that second acid molecule, to the action of a dehydrogenator, which eliminates one of its hydrogens.



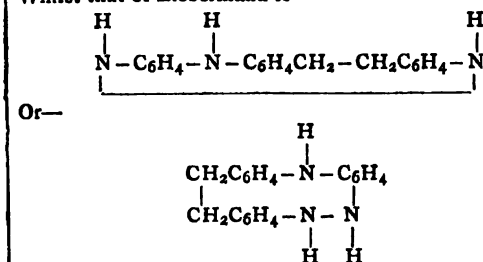
The group H—O— coming forth in the last reaction acts, as described above, on one of the hydrogens of the nitrogen appendix joined with the acid, and eliminates this hydrogen.

Here, as in the formation of violaniline and mauvaniline, both the nitrogens combine by their free atomicities, and form thus the closed chain representing the structural formula of rosaniline.

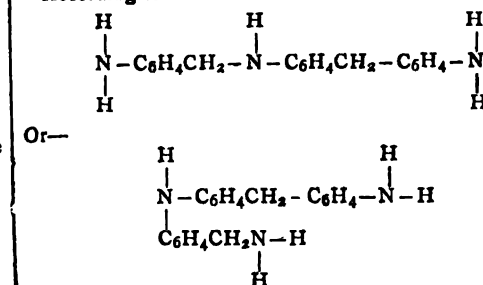
In this formula we see that a toluene nucleus is combined directly with a phenylene nucleus, whilst in Liebermann's formula two toluene nuclei are joined together, which is inconsistent with the mode of the formation of rosaniline. Liebermann's formula would have to be based on the supposition that the higher boiling toluidine is dissociated or acted upon before the much lower boiling aniline, which contradicts experience, and is improbable. Therefore we ought to give the preference to the formula being in conformity with the formation of rosaniline. This formula is—



Whilst that of Liebermann is—

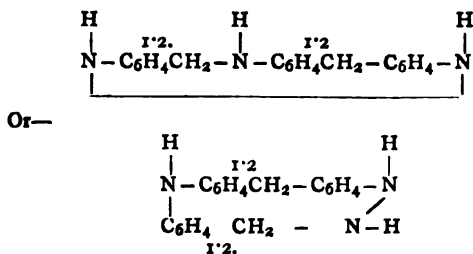


According to our formula leucaniline would be—



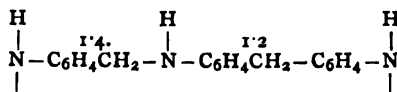
All reactions and decompositions of rosaniline and its salts can be explained as well by this new formula as by that of Liebermann. According to the different isomeric toluidines contained in the commercial aniline for red different isomeric rosanilines will exist.

Rosenstiehl's pseudo-rostaniline, made out of aniline and ortho-toluidine, would be represented by the following formula:—

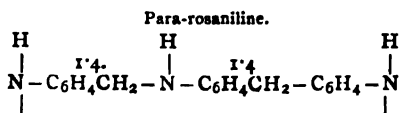


The commercial aniline for red contains, besides aniline, ortho- and para-toluidine, according to Beilstein and Kuhlberg and Rosenstiehl, both of which will contribute to the formation of rosaniline.

The boiling-point of ortho-toluidine being lower (197°) than that of para-toluidine (198°), the former will be acted upon the first, and therefore joined to the phenyl nucleus, and thus the formula of normal rosaniline will be—



According to Rosenstiehl (*Bull. Soc. Chim.*, [2], xi., 267) commercial magenta contains, in addition to rosaniline, an isomer of it, which he calls para-rostaniline, as it is formed entirely of aniline and para-toluidine.



There is a possibility of the existence of other rosaniline isomers than those named above, as perhaps those derived from meta-toluidine and aniline alone, or combined with the ortho- or para-toluidine, but they are not known at present. As all the aniline in the commercial aniline for red by the rosaniline process is used up for the formation of colouring matters, only toluidine salts remain, causing the formation of chryso-toluidines in a way similar to that of rosaniline, &c., as soon as the boiling-points of the respective toluidines are reached.

The boiling-point of aniline being 182°, that of toluidine say 198°, then the boiling-point of a mixture of two molecules of aniline and one molecule of toluidine is 188°, and that of a mixture of one molecule of aniline and two molecules of toluidine is 193°. The greater the difference between the boiling-points of such mixture the more decomposition takes place between these boiling-points, and therefore the quantities of the colouring matters produced are proportionate to the respective differences of boiling-points in a judiciously chosen mixture.

Boiling-point of aniline = 182°; of two anilines and one toluidine = 188°; difference = 6.

Boiling-point of one aniline and two toluidines = 193°; difference = 5.

Boiling-point of toluidine = 198°; difference = 5.

* The three different toluidines have three different boiling-points; 198° may be considered to be the average.

If a mixture of 10.5 parts of aniline and 5.5 parts of toluidine would be completely converted into colouring matters by the rosaniline process, a little less than 6 parts of violaniline, 5 parts of mauvaniline, and 5 parts of rosaniline are coming forth. If the proportion of aniline would be greater, a larger quantity of violaniline would be produced, and the yield of rosaniline would then be the less in proportion. If the proportion of toluidine would be greater larger quantities of chryso-toluidines would be produced, lessening the proportions of rosaniline too.

Practically, this process never takes place without formation of chryso-toluidines, the last being produced nearly simultaneously with the last produced quantities of rosaniline.

In the rosaniline process, practically the above-given quantities of aniline are converted completely in the above-given proportions of colouring matters; and thus, if we allow 2 parts more of toluidine for the formation of chryso-aniline in addition to the 5½ parts of toluidine, we see that 18 parts of a mixture of 10.5 parts of aniline and 7.5 parts of toluidine will yield nearly 6 parts of violaniline, 5 of mauvaniline, 5 of rosaniline, and 2 of chryso-toluidines, which is the greatest yield of pure and dry rosaniline crystals ever reached in practice.

It is evident that this yield cannot be increased at all, as we cannot alter the boiling-points of aniline and those of toluidines. The nearer the boiling-points are to one another, the more of the mauvaniline and rosaniline would be produced, and the less of violaniline and chryso-toluidines.

100 parts of a mixture of about 60 parts of aniline and 40 of toluidine are converted through the rosaniline process into colouring matters yielding 28 per cent pure and dry rosaniline, which represents 33.6 per cent of magenta crystals, with 7 per cent moisture. This would represent the largest yield of pure magenta crystals which could be reached. The greatest yield ever reached was not quite 33 per cent with arsenic acid, and 32 per cent of pure magenta crystals can be considered as a very good production; whilst with nitro-benzene never a yield over 30 per cent of pure magenta crystals has been attained, the used quantity of nitro-benzene calculated as aniline.

To find the best proportions of aniline and toluidine for the manufacture of magenta, we have to take the above given proportions and add to them an allowance for the quantities of aniline and toluidine which distil over in the rosaniline process, and are recovered also by steaming off the melt. This is a matter of practical experience, and varies according to the size of the charges and other circumstances.

Coupiér affirms that by treating his toluidine, which he declares to be a mixture of meta-toluidine (the latter is actually ortho-toluidine) and para-toluidine, with arsenic acid and hydrochloric acid he obtains from 40 to 50 per cent of rosa-toluidine, which is a confirmation of the above-given sentence, that the nearer the boiling-points the higher the yield of rosaniline, or, in that case, rosa-toluidine.

This thorough examination of the rosaniline process has led us to a more probable structural formula of rosaniline, to the explanation of the fact that the yield of rosaniline can never be higher than 30 per cent from 100 aniline for red, and to the respective positions of the nitrogen and methylene appendices in the three isomeric rosanilines known at present.

Adulteration of Geranium Oils.—M. Jaillard.—The author detects fatty oils, gum resins, and other liquid hydrocarbons as follows:—Into a test-glass are poured 5 c.c. alcohol at 70 per cent, and 6 drops of the oil in question, and the whole is well shaken up. If the oil is pure it remains bright and clear, while sophisticated specimens turn milky. This process is of course not available for the detection of cheaper ethereal oils.—*Wochenschrift Oel und Fett Handel.*

PROCEEDINGS OF SOCIETIES.

MANCHESTER LITERARY AND PHILOSOPHICAL SOCIETY.

Ordinary Meeting, April 15, 1879.

J. P. JOULE, D.C.L., LL.D., F.R.S., &c., President, in the Chair.

"Mineralogical Notes," by CHARLES A. BURGHARDT, Ph.D., The Owens College.

Precious Garnet (Almandine) from Ramsbottom, Lancashire.

A few days ago I received a small specimen of rock from a former pupil of mine (Mr. J. F. C. Sieber), accompanied by a note, in which the writer gave particulars respecting the locality in which the rock was found, and asked me to examine certain red granules present in the rock in order to ascertain whether they were garnets or not. The rock itself is a conglomerate of milky quartz grains (varying considerably in size), cemented together with silica and calcium carbonate; and disseminated throughout this conglomerate are the garnet grains in question, the whole constituting a 14-feet "fault" in the coal seam known as the Sand-rock or Featheredge Seam in the "Shipperbottom Mine," near Grant's tower at Ramsbottom. The coal from this mine is accompanied by a shale, which contains a very considerable amount of sulphate of aluminium, as it effloresces on being exposed to the action of the air; becoming eventually perfectly white like a piece of chalk. A microscopical examination of the rock showed the garnets to be very irregular in form and to vary considerably in size, the largest attaining a width of about 2.25 m.m., and the smallest about 0.75 m.m. They exhibited no distinct crystalline form, but a peculiar indented appearance was observed on the surfaces of some individuals, whilst others were pebble-like, somewhat convex, and rounded off as though they had been mechanically acted upon by water. The quartz grains did not exhibit the slightest trace of crystalline form, but were rounded off, and pebble-like. Both the garnets and the quartz grains could be extracted from the matrix, a cast of their forms being left behind. The red grains fused easily and quietly before the blowpipe to a black bead, which was not, however, magnetic: they were somewhat attacked by hydrochloric acid, with a slight separation of powdery silica; the presence of iron was also detected. The hardness of the mineral was 7.5, and its specific gravity 4.09 at 9° C. I hope shortly to determine the chemical composition. A microscopical examination of the garnet grains showed them to be crystalline, exhibiting certain marked peculiarities, and after a careful examination of many of them I came to the conclusion that the form in which they crystallised was the rhombic dodecahedron. I had previously proved them to be crystals of the regular system by examining the grains in polarised light and finding them to be isotropic. The crystals were built up out of myriads of minute laminated abnormal rhombic dodecahedrons, two faces of which have been enormously developed at the expense of the other faces. These laminæ are arranged parallel with each other in positions corresponding to each face of a rhombic dodecahedron, the indentations observed upon the indistinct crystal faces of some individuals being caused by this peculiar laminated growth. The laminæ vary from 0.0282 m.m. to 0.1880 m.m. in length, from 0.0188 m.m. to 0.1504 m.m. in width, and from 0.0031 m.m. to 0.0045 m.m. in thickness. The colour of the garnets (for such the crystals undoubtedly are) is light rose-red, and coupling this with their transparency and chemical properties there can be no doubt that they are alumina-iron garnets, or *almandine*. On examining some of the laminæ under the microscope I observed that they contain enclosures, which were mostly fluid cavities or "gas-pores," also here and there cracks and rifts, filled with a white substance, which most probably was calcite. On some of the laminæ there

was a separation out of a greenish mossy substance, which I believe to be ferric oxide, and on one individual I observed a few specks of iron pyrites. It is somewhat difficult to draw a conclusion as to the origin of these garnets in such a locality. From the microscopical examination of the rock it would appear that the cement and the garnets were both in a soft and pasty condition, the latter being probably formed whilst in the pasty matrix and prevented from attaining a normal development. It is a well-known fact that crystals which have formed in a free space or in a liquid mass, which has not exerted any retarding influence upon the force of crystallisation, always exhibit well defined faces and angles, whilst the opposite conditions result in abnormal growths such as the garnet crystals just referred to. The quartz grains were probably formed previously to the garnets, as they appear to be waterworn pebbles, and were most likely carried along with the semipaste like mass which contained the constituents of garnet and calcite in its substance. When the paste hardened, the quartz grains and the garnets were enclosed in the manner exhibited on the specimen of rock from Ramsbottom. All the constituents of garnet were at hand, namely, aluminous shale, which would furnish the alumina and silica, and iron pyrites, which would furnish the iron, whilst water impregnated with calcium carbonate, acting upon the shale and iron pyrites, would bring about the necessary chemical changes, calcium sulphate being formed and carried away, and the garnet solution (if such a term may be used) could then crystallise out. From the formation and the locality in which the garnets were found it is quite certain that their origin was aqueous. Another explanation of the occurrence of garnet in the conglomerate may be as follows, viz.:—Garnet is a very common accessory of crystalline rocks, occurring mostly in slates of various kinds, such as talc slate, mica slate, chloritic slate, and aluminous slate; it also occurs in gneiss, granite, porphyry, serpentine, and granular limestone. One or more of the above-mentioned rocks containing crystallised garnets already formed were disintegrated by the action of water, and the resulting *débris* (consisting of grains or pebbles of quartz and garnets) was carried away and deposited in a semi-fluid mass of silica and calcite, and thus cemented together. The question which next arises is, where and what is the original garnet rock in the neighbourhood of Ramsbottom? Is it perhaps the aluminous shale? This problem still remains to be worked out. It is evident, however, at once on examining the conglomerate that there is a large excess of quartz grains present, with a very fair sprinkling of garnets; whilst the cement is not present in any great amount. An examination of thin sections of this rock under the microscope may clear up some doubts and throw further light upon its origin.

Fibrous Rock-salt (Sodium Chloride).

This anomalous formation has always been an interesting one on account of the strong cubical "habit" of rock-salt, all other forms than that of a well-defined cube being rarities. The fibrous formation is not due to a crystallisation of the sodium chloride in another system than the regular system, but is explained by an abnormal development of four cubical faces parallel to one axis, which may with propriety be called the prismatic axis. So far as I am aware, the formation of fibrous rock-salt has not been explained. Having the well-known fact before me, viz., "the influence exerted upon the crystal form of a body crystallising out of its solution by the presence in that solution of a small quantity of a foreign body," I thought it probable that a similar effect might be produced upon the growth of sodium chloride crystals by the presence of a foreign body or bodies. In order to ascertain this I made a strong saturated solution of ordinary table-salt, filtered it, and precipitated out most of the salt by passing hydrochloric acid gas into the solution. The liquid was drained off from the precipitated salt, and allowed to evaporate *very slowly* for several days in a narrow necked flask, and then laid

aside to cool. I then observed amongst the mass of ordinary salt numerous long well-developed prisms. Some of these prisms were an inch or more in length, and sometimes terminated by a broad cube. Others, again, were twinned according to the usual law, viz., "the twin plane a face of the octahedron." I observed that many of the cubical crystals were somewhat bent, and resembled closely the bent barytes crystals of the prism and basal terminal plane in combination. Nearly all the crystals (prismatic and cubical) were opaque or almost so, and of a very white colour. In addition to free hydrochloric acid, I found in the solution a quantity of magnesium sulphate and magnesium chloride, and it is to the peculiar retarding action of these substances, either together or singly, that I ascribe the general formation of fibrous rock-salt; for they could easily occur *naturally* in any rock-salt formation. As sulphate of magnesium and chloride of magnesium are constituents of sea-water, and generally present in all waters flowing from rocks, either crystalline or sedimentary, their presence during the deposition of rock-salt is ensured, whilst the presence of hydrochloric acid would also be possible if a slight rise of temperature took place during the deposition, it being a fact that magnesium chloride gives off hydrochloric acid during evaporation at comparatively low temperatures.

In the discussion which followed the reading of the above paper Mr. BINNEY, F.R.S., read an extract from *Trans. Man. Geol. Soc.*, vol. i., p. 87, to the following effect, viz.:

"On the Marine Shells found in the Lancashire Coal Fields," by E. W. BINNEY.

"Immediately above the 'Rough rock' at Birtle Dean, near Heywood, is the Featheredge coal, about two feet thick; this seam has a roof of one yard in thickness, consisting of black shale which is full of the *Pecten*, *Goniatites*, *Povidonia*, &c., mixed with various species of *Ferns*, *Lepidodendra*, and *Sigillariæ*. It is a singular fact that the black shale has scarcely ever been seen on the roof of this coal, except at Messrs. Ramsbottom's Colliery, at Birtle Dean. At Fecit, within two miles of the last named place, and other localities, the roof of the coal consists of a coarse conglomerate of quartz pebbles, containing numerous garnets and crystals of carbonate of barytes, generally presenting a water-worn appearance. I have also observed small pieces of opal imbedded in this rock."

NOTICES OF BOOKS.

Science Teachings in Living Nature. A Popular Introduction to the Study of Physiological Chemistry and Sanitary Science. By W. H. WATSON, F.C.S., F.M.S. London: E. Stanford.

THIS book labours under the disadvantage of being, to a very great extent, an *Itas post Homerum*. Had it appeared prior to such well-known and popular treatises as Johnston's "Chemistry of Daily Life," Lewes's "Physiology of Daily Life," and others which might be mentioned, it would have achieved and deserved a great success. As the case actually stands, however, containing little either in facts or in conclusions which may not be found in the works just referred to, we fear it will be regarded as anticipated.

Although the author's teachings are upon the whole in accordance with what is regarded as demonstrated, there are a few cases where we find statements either questionable in themselves or capable of being misunderstood by the unscientific reader, for whom the book is avowedly compiled.

Thus in the Introduction (p. x.) we are told that "the brains of the wise . . . are explained by the necessary application of chemical science." This sentence may convey the impression that chemistry has succeeded in

throwing a light upon the connection between intelligence and that part which we assume to be its organ, and in discovering a difference in composition or molecular arrangement between the brain of the wise and of the unwise. Mr. Watson must be well aware that no such explanation has been given, and that the manner in which the brain subserves the action of the reasoning principle within us is scarcely less a mystery to us than it was to our rudest forefathers.

On page 9 the author contends—as was at one time very generally believed, and as is still held by General Pleasonton and his followers—that "the value of light in relation to life depends, up to a certain point, upon the amount of the blue and violet rays which it contains." This view is not in harmony with the results of more recent observations, nor does it agree with certain conclusions quoted by the author on a subsequent page.

In the outset of Chapter II. Mr. Watson seems to adhere to the old supposition, now abandoned, that the respiratory process in plants and in animals is essentially different and opposite. Thus he writes:—"Fungi are a peculiar exception to this rule, for their breathing functions resemble those of animals rather than those of plants." It is now known—see, e.g., Prof. Allman's Presidential Address at Sheffield—that the respiration, strictly so-called, of animals and plants is identical, oxygen being in each case absorbed and carbonic acid evolved. In plants, however, during daylight, the respiratory process is masked by the much more active process of nutrition, in which carbonic acid is decomposed, its carbon assimilated, and its oxygen liberated. Mr. Watson is doubtless aware of this distinction between vegetable respiration and nutrition, and of the identity of the breathing process in the two great organic kingdoms; but we fear that the reader will fail to find it in his pages.

On page 67 we read—"It has been well said that a bad theory is worse than none at all." With this dictum we can agree only to a very limited extent. Even a bad theory gives a definite character, which might otherwise be wanting. The mischief begins when men ignore or garble facts which cannot be made to chime in with their theories. The doctrine of phlogiston—which, by the way, was not held in ancient times—was useful in the earlier part of its existence.

Turning over the leaf we come upon a passage which we must beg to quote:—"Lavoisier, whose history is perhaps the most noteworthy of any philosopher either before or since, was put to death owing to the scientific opinions which he held; and looking through the histories of science generally, I think we cannot fail to notice that religion has from very early times been a damper to the progress of science."

We certainly never before heard it insinuated that any of Lavoisier's scientific opinions were objected to by the heads of the Revolution. The charge on which he was arraigned and condemned was that, in his capacity of *Fermier-Général*, he had connived at the adulteration of certain tobacco. The judges certainly declared that "the Republic had no need of chemists," but this by no means bears out Mr. Watson's assertion. However this may be, anyone reading the sentence we have quoted can scarcely avoid inferring that Lavoisier's persecutors were ecclesiastics, or were actuated by a religious motive, whilst in point of fact they were mainly atheists! We admit that "scientific savans," as Mr. Watson calls them, have been oppressed by "clericals." But we think the future enemies of Science must be sought elsewhere. Let it once be understood that she has no word of cheer for those who would thrust God out of the universe, and that her teachings—contrary to the vain fears of Virchow—are hostile to "social democracy," she will find enemies who merely wait their opportunity.

Towards the end of the work we are told that "most of our large towns are now supplied [with water] from the Lake Districts." This assertion deals probably with the future rather than the present.

We should suggest to Mr. Watson a reconsideration of the passages to which we have called attention.

Lectures on the Theory and Control of Infectious Diseases.

By JAS. B. RUSSELL, M.D., Medical Officer of Health. And on *Air, Water Supply, Sewage Disposal, and Food.* By W. WALLACE, Ph.D., F.C.S., City Analyst. Glasgow: Maclehose. London: Hamilton, Adams, and Co.

It has often been remarked that the number of chemical manuals produced in this country is out of all fair proportion to our yield of chemical research. In like manner it might be said that the progress of rational measures for the promotion of public health has been scarcely in accordance with the number of sanitary treatises, lectures, and speeches with which we are favoured. Indeed—save among those whose official duties compel them to labour, in season and out of season, at the repression of infectious disease, and, on the other hand, among the more favoured few to whom sanitary movements have brought emolument and celebrity—the interest in the subject is perceptibly declining. Ten years ago, for instance, the disposal of sewage was often eagerly and intelligently discussed among persons casually meeting, *e.g.*, in public conveyances. This excitement has died out, and the average John Bull is perhaps a little too ready to pronounce the various systems of sewage-treatment to be mere costly humbugs; sport, possibly, to engineers, but death to the rate-payer. That the lectures before us contain a great amount of interesting matter, the names of its authors will be a sufficient guarantee. Of the many passages which might claim especial comment, space will permit the notice of a few only. Thus in a quotation from Dr. Farr, with which Dr. Russell's first lecture opens, we find it mentioned that "they"—*i.e.*, infectious diseases—"take the lives of criminals that justice has not condemned." Surely in so far these diseases render a great service to society, and save justice much trouble. On the important subject of density of population as affecting mortality, we find the persons per square mile in the Manchester District given as 22,357, and in the Liverpool District 65,823. Yet this fivefold increase of the density of population merely raises the death-rate from 38 per thousand in the former case to 39 in the latter.

Dr. Alison's painful story of the spread of typhus fever, quaintly presented by Thomas Carlyle,* is no doubt a striking reply to those who look with apathy upon disease among the poor. But it has yet a further lesson: if community of disease proves kindred, how can we still maintain the old myth of the total and essential distinction between man and the lower animals?

Dr. Russell contends, and in our opinion with perfect justice, that poor-law administrators and poor-law officials are essentially unfitted for the successful performance of the functions of a local authority under the Public Health Act.

The importance of vaccination is ably pleaded, but the difficult question still remains unsolved: how is it that now this supposed safeguard is all but universal, we have still small-pox epidemics, whilst in the earlier part of the century (say about 1830), when vaccination was optional, and was rare save among the more cultivated classes, the disease seemed to have vanished, and was spoken of as a thing of the past?

Dr. Wallace, in his comparison between the respective production of smoke in Glasgow and in London, overlooks the great distinction that Glasgow is essentially a manufacturing town, whilst London carries on manufactures only to a very subsidiary extent. A law empowering the magistrates in Glasgow, Leeds, &c., to order the closing of works which cannot be conducted without the emission of smoke would be, in our opinion, a most rash and impolitic measure. The lecturer complains that "avarice" entails upon us "narrow streets, built up lanes, and every foot of

ground converted into money." But if we insist upon wider streets the same avarice will charge proportionately higher ground-rents.

We cannot admit that "both the mechanical and dissolved impurities of water" are excluded on freezing. We have repeatedly seen ice in which sticks, straws, vegetable fibre, and particles of excrementitious matter have become imbedded.

We are very glad to see a protest against the sensational system of describing all matters present in water, save oxygen and hydrogen, summed up as "total solid impurity." It misleads the public. From the abstract chemical point of view small quantities of carbonate of lime, &c., may doubtless be regarded as impurities. But when we are considering a drinking water, from a dietetic point of view, the case is different. That absolute H₂O would be a salutary beverage is a totally unproved assumption.

The author's remarks on the various systems of sewage-treatment are not reassuring. Without entering upon a controversy, we will merely express the wish that he would have considered some of the evidence by which certain of the conclusions of the Rivers' Pollution Commissioners have been so superabundantly refuted.

In the interesting lecture on food Dr. Wallace remarks that the "only materials" used in baking bread are flour, water, yeast, and salt. In London he would be told by anyone connected with the trade, that he had forgotten that legalised adulteration, the potatoes.

We trust that these courses of lectures may have had a beneficial effect, as we have heard it contended in Glasgow—and that by one of its citizens of no mean standing—that all the impurities of the Clyde and the Kelvin, and all the contaminations of the air, had wrought no harm to the health or the longevity of the inhabitants.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 11, September 15, 1879.

On Woollen Cloths Dyed a Blackish Blue in order to Replace the Indigo-blue used in the Uniforms of the French Army.—E. Chevreul.—The author shows that the colouring-matter of the cloth in question cannot be indigotin, Prussian blue, or ultramarine, but that it may be one of the aniline dyes.

Experiments tending to show the Compound Nature of Phosphorus.—Letter from Mr. N. Lockyer to M. Dumas.—Phosphorus heated in a tube with copper gives a very brilliant hydrogen spectrum. Phosphorus alone, heated in a tube where a vacuum has been formed by Sprengel's apparatus, gives nothing. Phosphorus, at the negative pole in a similar tube, gives very abundantly a gas which shows the spectrum of hydrogen, but which is not hydrogen phosphide.

Determination of Organic Nitrogen in Natural Waters.—H. Pellet.—For the determination of ammoniacal nitrogen the author follows the directions of Boussingault. For the nitric nitrogen he evaporates 3 litres of water, and when the residue amounts merely to 60 or 80 c.c., he adds acetic acid to decompose the carbonates without attacking the nitrates. The whole is heated to a boil and the volume is made up to 100 or 200 c.c., according to the deposit. He then filters and determines the nitric nitrogen in 25 or 50 c.c. of the liquid according to Schloesing's method, measuring the binoxide of nitrogen produced. For the total nitrogen he likewise

* "Past and Present," Book III., Chap. .

evaporates 3 litres of water in contact with 2 grms. of pure magnesia to drive off the ammoniacal nitrogen. A part of the dry residue is mixed with soda-starch and introduced into a combustion-tube. By this addition of carbonaceous and hydrogenous matter all the nitric nitrogen passes into the state of ammonia, and the operation becomes an ordinary determination of nitrogen by the soda-lime process. The quantity of nitrates present, however, should not be greater than 0.20 or 0.25 gm. of nitrate of potassa. This is the reason for the previous separate determination of the nitric nitrogen.

Oxidising Action of Cupric Oxide: Transformation of Acetic Acid into Glycolic Acid.—P. Cazeneuve.—10 grms. acetate of copper, powdered and mixed with 25 grms. of water, were heated for an hour to 200° in a sealed tube. The tube contains a deposit of cuprous oxide in crystals, apparently of a special form. On opening the tube a little carbonic acid escaped, and the filtrate, on evaporation in a vacuum at a temperature not exceeding 50°, deposited blue crystals, which prove to be glycolate of copper. By an analogous reaction the propionic acid may be transformed into the lactic.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin,
No. 8, 1879.

Oxidation of Cinchonin-chinolin by Potassium Permanganate.—W. Koenigs.—The author is attempting to break up the benzol ring of chinolin by the oxidation of this base or of amido-chinolin, in the hope of thus forming a phthalic acid of pyridin.

Piperidin and Pyridin.—A. W. Hofmann.—The author has examined the action of bromine upon piperidin, obtaining a compound $C_5H_7Br_2NO$, with its platinum-silver and methyl combinations. Pyridin has likewise been treated with bromine, the result being dibrom-pyridin.

Angelyl Mustard Oil.—A. W. Hofmann.—The compound obtained, C_6H_9NS , is a higher homologue of crotonyl oil of mustard.

Umbelliferon and some of its Derivatives.—F. Tiemann and C. L. Reimer.—The authors prove that umbelliferon is an oxy-cumarin derived from resorcin.

Aldehyds from Orcin and their Derivatives.—F. Tiemann and E. Helkenberg.—The compounds described are orcyl-aldehyd, orcyl-aldehyd-anilid, homo-acetoxycumarin, α -orcendialdehyd, α -orcendialdehyd-dianilid, and β -orcendialdehyd.

Tribrom-phenol-brom and Tribrom-resorcin-brom.—R. Benedikt.—The former of these bodies is formed on the exhaustive bromation of many aromatic oxy-acids. The penta-brom-resorcin of Stenhouse and the tribrom-resoquinon-bromide of Liebermann is probably tribrom-resorcin-brom.

Fractionated Combustion of Hydrogen and Marsh-Gas.—W. Hempel.—A mixture of hydrogen with excess of oxygen, passed over palladium burns completely. The reaction begins at common temperatures, the palladium becomes red hot, and an explosion follows if the gases are present in a suitable proportion. Marsh-gas with oxygen, passed over palladium, burns at about 200°. Mixtures of hydrogen, marsh-gas, and oxygen, if in the proper proportion, detonate violently. In such mixtures, when the oxygen is in excess, the hydrogen alone burns if the external temperature does not exceed the marsh-gas remaining unattacked, provided the palladium is prevented from becoming too hot.

On Nitro-alizarin.—E. Schunck and H. Roemer.—The authors find that their method of nutrition (*Berichte*, xii., p. 384) had been already made known by Caro.

A Homologue of Phosphenyl-chloride.—A. Michaelis.—Along with phosphenyl-chloride the author has obtained a homologue of the composition $C_7H_7PCl_2$.

Reply to the Remarks of E. and O. Fischer.—O. Döbner.—A continuation of the "malachite green" controversy.

Biedermann's Central-blatt für Agrikultur-Chemie.
September, 1879.

Researches on the Proportion of Carbonic Acid in the Atmosphere.—J. Reiset.—Taken from the *Comptes Rendus* and already noticed.

Investigations on Spongy Iron and Animal Charcoal as Agents for the Purification of Water.—Dr. L. Lewin.—A perfect filtering material should be able to remove not merely suspended bodies, but such as are in states of physical and chemical combination, and either retain them in such a manner that the subsequent application of pure, or at least purer, waters may be unable to wash them out, or else to transform them into a condition in which they can no longer interfere with the quality of the water. In addition a filter must retain these powers unimpaired for a length of time. The attempts which have been made to imitate the processes of nature by passing the river water used for the supply of large towns through deep beds of sand and gravel have shown that such a process has no claim to be regarded as chemical. The case is different when charcoal, and especially animal charcoal, is used as filtering material. The latter not only holds back colouring-matters and decomposes or partially retains most mineral salts and gases, but it absorbs and detains the most varied organic compounds, nitrogenous as well as non-nitrogenous. Particularly important is its power—first demonstrated by Claude Bernard, and confirmed by other inquirers—of absorbing albumen, a power which increases with the quantity of the latter, with the time of contact, and with the concentration of the liquid to be filtered, and which is the more important as the arrested body is held so firmly as not to be removed by solvents. This valuable material has also its shortcomings, which especially consist in its ready saturation with impurities so that it needs to be revived, i.e., freed from the detained matters, or if this is impracticable, to be renewed altogether. Other substances have therefore been suggested as substitutes for charcoal, and amongst others various modifications of iron, especially spongy iron. The latter substance has been submitted by the author to an elaborate experimental examination. His experiments show that as regards nitrogenous substances the spongy iron filter is inactive. Such substances are neither decomposed nor held back in such a manner as not to be readily washed out again. It can be stated with the fullest certainty that if only 10 c.c. of urine containing moving bacteria are mixed with 2 litres of water, and passed through the spongy iron filter, numerous bacteria can be detected in the filtrate. The odour of putrescent matter is but very slightly removed by spongy iron. The alleged improvement undergone by plumbiferous waters in contact with spongy iron were found to be insignificant.—*Zeitschrift für Biologie*, 1878, No. 4, pp. 483—506.

A Contribution to the Chemistry of Starch.—Dr. V. Griessmayer.—The following transformation products originate on the action of diastase or of dilute sulphuric acid upon starch:—(1.) Soluble starch, insoluble in water of from 50° to 60°; in the solid state it is coloured by iodine blue, but in an aqueous solution a vinous red, and if dried up with an excess of iodine it becomes violet, yellow, or brown. (2.) Erythro-dextrin, which is chiefly met with in commercial dextrin; it is never insoluble in water, and is coloured red by iodine, both in the solid state and in solution. (3.) The achroo-dextrins, α , β , and γ , which take no colour with iodine. α can be partially converted into sugar, but less easily than 1 and 2. β resists the action of diastase for twenty-four hours at least. γ is not attacked by diastase for a year at least. (4.) Maltose, not readily attacked by diastase. (5.) Glucose.

The author considers starch as a poly saccharide of the formula $n(C_{12}H_{20}O_{10})$, in which n must be first determined, but cannot be less than 5 or 6.

La Correspondance Scientifique.
September, 1879.

This issue contains the presidential discourse of M. Bardoux, delivered at the Montpellier meeting of the French Association for the Advancement of Science. Among the papers read before the Chemical and Physical Sections were:—On the Compound Ammonias, by M. Buisine; on the Oxidising Action of Cupric Oxide, by M. Cazeneuve; on the Determination of Methyl Alcohol in Ethylic Alcohol, by M. Caillol; on the Calcareous Soils of Barcelona from a Chemical Point of View, by M. Luis Cabello e Ibanez; on the Emissive and Absorbent Powers of Flames, by M. Rosetti; on a Compass for the Measurement of Powerful Currents, by M. Ducrétet; Method of Measuring the Intensities of the Currents of Bunsen Batteries by Means of Thomson's Galvanometer, by M. Mercadier; and on Thermometric Measurements of High Temperatures, by M. Crova. The "compass" of M. Ducrétet is a modification of the well-known instrument of Pouillet. The vertical circle receives a different arrangement, and instead of being fixed perpendicularly it is mounted at its centre by a system of pivots, which traverse it in the direction of its diameter. Hence it becomes possible to give it all positions comprised between the vertical and the horizontal.

Chemiker Zeitung.

No. 33, August 14, 1879.

The German Imperial Public Health Department is said to be preparing measures for the repression of secret and proprietary remedies.

Determination of Sugar in Beet-root.—Dr. Scheibler.—It has been falsely assumed that all the liquid in the root contains water. The author shows that in addition to the saccharine sap there exists in the beet non-saccharine water, chemically combined with the solids.—*Arch. Pharm.*

No. 34, 1879.

In Switzerland the sale of patent and secret medicines is about to be submitted to very stringent regulations.

Lute for Distillatory Apparatus.—Thanisch, instead of linseed-meal, proposes strips of brown paper smeared over with book-binders' paste to which one-eighth of glycerin has been added.

No. 35, 1879.

Chemical Purification of Waste Waters.—Jean de Molins.—The author, who has dealt with the waste waters from the woollen mills of Roubaix, recommends milk of lime and sulphate of alumina, together or separately. He considers that the action of the aluminous hydrate upon the organic impurities of water is quite analogous to its behaviour with dissolved colouring-matters which it throws down in the form of lakes. He also points out that clay, if diffused in water, becomes coagulated by the presence of certain salts, and carries the organic impurities down with it.

New Method of Producing Varnish.—Dr. E. Schrader.—The author causes ozone to act upon linseed oil, which is at once perfectly bleached and brought to the proper consistence without the aid of fire.

Quantitative Determination of Tannin.—Dr. Ostermayer.—The author proposes the following modification of Wagner's process. 10 grms. of the substance are completely exhausted with hot water; the liquid filtered when cold, and evaporated to dryness. The residue is extracted with a mixture of 1 part ether and 2 parts alcohol at

90 per cent as long as tannin is taken up. The extracts are united, again evaporated to dryness, and the residue dissolved in 500 c.c. water. The solution is almost colourless. 50 c.c. of this solution are titrated with a solution of cinchonin coloured with magenta, till the liquid begins to retain the colour of the latter. Every c.c. of cinchonin solution consumed corresponds to 1 per cent of tannin. To prepare the cinchonin solution 4.523 grms. of cinchonin sulphate, 0.1 grm. magenta, and and 0.05 grm. sulphuric acid are dissolved in water and made up to 1 litre.—*Pharm. Zeitung.*

Iodine-starch Reaction.—E. Heintz.—The author shows that tannin is among the substances which can prevent the appearance of this reaction.—*Pharm. Zeitung.*

Reaction for Salicylic Acid.—H. Schulz.—If an aqueous solution of salicylic acid or sodium salicylate is mixed with a little solution of copper sulphate a bright emerald green colour appears, perceptible if 1 part of the sodium salt is dissolved in 2000 parts of water. The addition of a stronger acid, or of ammonia, destroys the green colour.—*Pharm. Zeitung.*

Distribution of Arsenic in the Organism after Ingestion of Arsenious Acid.—Prof. E. Ludwig, after the examination of many cases, both of acute and chronic poisoning, finds the arsenic chiefly collected in the liver, very little being retained in the brain and the bones. This conclusion is directly opposite to that of Scodossoff, who always found most arsenic in the brain.—*Akad. Wissensch. Wien.*

Occurrence of Algæ in Saline Solutions.—A. Tschirch makes the interesting observation that in almost all bottles of magnesium sulphate, and in some of calcium sulphate solutions, in the laboratory of Berlin University, algæ have made their appearance. They are Palmellaceæ and develop chlorophyll.

No. 36, 1879.

This issue opens with a short biographical notice of Berzelius, the centenary of whose birthday was celebrated on August 29.

A serious explosion took place on August 15 in an oil and colour warehouse in Vienna, in consequence of a carboy of benzin being decanted in the proximity of a naked flame. The proprietor and three assistants are dead from the wounds received.

The Alps and the Jura are being diligently searched by gold hunters. Certain English speculators have purchased and re-opened a mine in the Grisons, locally known as "Die Goldene Sonne."

Dr. Brauns, of Halle, has been appointed to the chair of mineralogy and palæontology in the University of Tokio, Japan.

MISCELLANEOUS.

University College, London.—A special Practical Course of instruction has been arranged to meet the requirements of Students who are preparing for State Medicine Examinations, or who wish to become Public Analysts. It will consist of two parts, and will require daily work for four months. It may be commenced at any time during the Session, from October to July. The State Medicine Department is under the direction of Professor Corfield, M.A., M.D.; Mr. C. E. Cassal, F.C.S. being demonstrator in the Hygienic Laboratory.

University College, Bristol.—In addition to the Courses of Lectures and Laboratory instruction referred to in our Students' Number arrangements have been made for a special Course, by Professor Letts, on Metallurgy, explaining the method of extraction, purification, &c., of the useful metals, and the properties upon which their utility depends. The fee for two terms is two guineas.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

SEPTEMBER, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

	Appearance in 2 foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	An- Sulphuric hydride.	Hardness on Clark's Scale		
		Saline.	Organic.								Before Boiling.	After Boiling.	
		Grs.	Grs.								Grs.	Degrees	
<i>Thames Water Companies.</i>													
Grand Junction	Clear	0'000	0'009	0'126	0'148	20'00	7'750	0'504	1'008	1'133	14'1	3'6	
West Middlesex	Clear	0'000	0'010	0'106	0'159	20'40	7'590	0'432	1'008	1'133	14'1	3'3	
Southwark and Vauxhall	Clear	0'000	0'009	0'105	0'155	20'20	7'160	0'460	1'080	1'330	13'2	3'7	
Chelsea	Clear	0'000	0'009	0'108	0'089	20'00	7'630	0'432	1'152	1'330	15'2	4'6	
Lambeth	Clear	0'000	0'011	0'020	0'148	20'70	7'340	0'496	1'080	1'630	14'7	3'3	
<i>Other Companies.</i>													
Kent	Clear	0'000	0'001	0'390	0'014	26'70	7'460	0'632	1'800	1'660	18'8	6'5	
New River	Clear	0'000	0'004	0'132	0'094	20'70	7'400	0'576	1'080	1'630	13'7	3'3	
East London	Clear	0'000	0'009	0'126	0'062	20'80	7'160	0'432	1'152	1'300	13'2	3'7	

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

TO CORRESPONDENTS.

J. A.—The books named give full information.
H. V.—Auerbach is probably right.

NOTES AND QUERIES.

Pyrites in Manufacture of Sulphuric Acid.—Could any of your readers inform me which is the cheapest way of using up smalls of pyrites in the manufacture of sulphuric acid?—“SULPHUR.”

Analysis of Cinchona Bark for Alkaloids.—Can any of your readers inform me what are the details of the process, or where the process may be found, for the complete analysis of cinchona bark for alkaloids as followed by London quinologists?—L.

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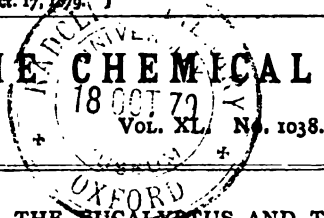


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THE CHEMICAL NEWS.



THE EUCALYPTUS AND THE PINE
CONSIDERED IN RELATION TO THEIR
SANITARY PROPERTIES.*

By C. T. KINGZETT, F.C.S., F.I.C.

As is widely known, both the eucalyptus and the pine have long enjoyed a popular reputation as health agents. In particular the eucalyptus has acquired the character of the "fever-destroying tree," and has been largely cultivated in various parts of the world with the view of rendering habitable large districts previously devastated and depopulated by malaria, &c. In this paper, the author, referring to the old theory of the action of the eucalyptus, which attributes its sanitary powers to its drainage properties, shows that this hypothesis always lacked evidence, and at its best was of a negative character. In the first place, the eucalyptus is only superior to other trees as a means of drainage in the proportion in which its rate of growth exceeds that of other trees, and this is not sufficiently notable to account for its extraordinary fever-destroying properties. Then, again, other trees, even if planted in malarial districts do not free them from the disease, so that the action of the eucalyptus is of a positive type, and, like the pine tree, its properties are of a healthful nature, upon whatever soil or in whatever climate it may grow, whether in deep valleys or upon the sides of mountains. Others have maintained that just as pine forests are supposed to exert their beneficial influence upon persons suffering from pulmonary and other affections, by virtue of the volatile emanations arising from them, so the eucalyptus produces its well known effects by the oil which is evaporated from its leaves. Mr. Kingzett then examines this hypothesis in detail, and shows the relative oil-yielding power of the different species of eucalyptus. The genus embraces over 130 species, and of these *Eucalyptus amygdalina* is the most abundant oil-giving tree; 100 lbs. of the leaves giving from 3 to 6 lbs. of the oil. This oil is practically identical in composition with the oil of turpentine derived from pine trees, and with most of the so-called essential oils or perfumes. By the investigations of Mr. Kingzett, it has been ascertained that all these oils when subjected to the action of atmospheric oxygen and moisture produce peroxide of hydrogen and a number of camphoraceous substances having marked antiseptic powers. Knowing, then, how much of these substances are yielded in the laboratory by a given quantity of oil of eucalyptus or oil of turpentine, Mr. Kingzett has extended his calculations to the pine and eucalyptus forests, which are so abundantly distributed in nature. Taking New South Wales and South Australia alone, he calculates that the eucalyptus forests of this district contain at any given moment sufficient oil in the leaves (ready to be evaporated into the atmosphere under the agency of warm winds) to form by contact with the atmosphere no less than 92,785,023 tons of pure peroxide of hydrogen, and 507,587,945 tons of camphoraceous principles. Now, if it be remembered that in nature all matters of animal and vegetable origin are oxidised by the atmosphere, which is thus kept free from the pernicious products of putrefaction, and that peroxide of hydrogen is a much more powerful oxidant than ordinary oxygen, and if it be also borne in mind that the camphoraceous products above referred to are also powerful antiseptic

* Abstract of a paper read before the Health Section of the Social Science Congress, Manchester.

agents, then the healthful influences of the eucalyptus can neither be wondered at nor be longer open to any doubt. What is true of the eucalyptus is true also of the pine, and on an immensely larger scale, for pine forests are distributed freely over both hemispheres, and the oil of turpentine, which is a natural product of these trees, undergoes the same chemical changes in the atmosphere as oil of eucalyptus. By imitating this natural process of oil oxidation, Mr. Kingzett has, as is well known, succeeded in obtaining and rendering available in commerce the antiseptic and oxidising principles to which pine and eucalyptus forests owe their hygienic influences.

MENDELJEFF'S PERIODIC LAW AND THE
MAGNETIC PROPERTIES OF THE ELEMENTS

By THOMAS CARNELLEY, D.Sc.

IN 1845 Faraday showed that all matter was divisible into two classes—(1) *Para-magnetic substances*, or those which are attracted by a magnet and set axially when placed between its poles; (2) *Diamagnetic substances*, or those which are repelled by a magnet and set equatorially, or at right angles to the first position when placed in the magnetic field. He also determined qualitatively the magnetic properties of a large number of elements in the free state, with the following results ("Experimental Researches," vol. iii.) :—

Para-magnetic Elements.—N, O, Fe, Ni, Co, U (Verdet, *vide* Lloyd's "Treatise on Magnetism"), Mn, Pt, Os, Pd, Ir, Rh, Cr, Ti, Ce, C (Ganot's "Physics," p. 807), K (Lamy).

Diamagnetic Elements.—H, Na,* Cu, Ag, Au, Hg, Zn, Cd, Pb, Sn, P, As, Sb, Bi, S, Se (Becquerel), Cl, Br, I. Also Ti (Crookes) and Si (Lamy, *Ann. Chim. Phys.*, li., 305).†

On comparing together the elements contained in these two classes, we seek in vain for any evident connection between their magnetic and chemical properties. For in both classes we have metals and non-metals, and in this arrangement many elements are separated which otherwise are nearly related, as in the case of K and Na, O and S, Au and Pt, N and P, As, Sb, &c. It has been stated (Watts's "Dictionary of Chemistry," iii., 773; Lothar Meyer's "Modernen Theorien der Chemie," p. 233) that elements with small atomic volumes as Fe, Ni, Co are mostly para-magnetic, whilst those with large atomic volumes are mostly diamagnetic. But to this rule there are many exceptions: thus K, with large atomic volume, is para-magnetic; and H, Cu, Ag, Au, Zn, Hg, and Cd, with small atomic volumes, are diamagnetic.

In his well-known memoir on the Periodic Law (*Ann. der Chem. u. Pharm.*, Sup. viii., 133) Mendeljeff gives a classification of the elements similar to that represented in the table below, and he points out that those elements of a given group which belong to even or to odd series resemble one another much more nearly than the odd members resemble the even members of the same group. I have lately been working on the subject of the influence of atomic weight on the properties of elements and compounds, and it occurred to me to compare the above magnetic list with this classification of Mendeljeff. When this is done, a most remarkable confirmation is obtained of the above theory of odd and even members of a group of elements, such that the following rule *invariably* holds good without a single exception in the case of the thirty-eight elements to which it can be applied :—*Those elements belonging to the even series of Mendeljeff's classification*

* According to Faraday Na is strongly diamagnetic, whereas Lamy states it to be slightly magnetic.

† U and W in combination are, according to Faraday, very slightly diamagnetic; the magnetic properties of W in the free state have however, not been determined.

Series.	Group I.	Group II.	Group III.	Group IV.	Group V.	Group VI.	Group VII.	Group VIII.
1	-H=1							
2	? Li=7	? Be=9.2	? B=11	+C=12	+N=14	+O=16	? F=19	
3	-Na=23	? Mg=24	? Al=27.4	-Si=28	-P=31	-S=32	-Cl=35.5	+Fe=56; +Co=59;
4	+K=39.1	? Ca=40		+Ti=48	? V=51.3	+Cr=52.3	+Mn=55	+Ni=59
5	-Cu=63.5	-Zn=65.2	? Ga=69		-As=75	-Se=79	-Br=80	+Ru=104; +Rh=104;
6	? Rb=85.5	? Sr=87.5	? Yt=89	? Zr=90	? Nb=94	? Mo=96		+Pd=106
7	-Ag=108	-Cd=112	? In=113	-Sn=118	-Sb=122	? Te=128	-I=127	
8	? Cs=133	? Ba=137	? Di=138	+Ce=140				
9		? Ng=145*						
10			? Er=178	? La=180	? Ta=182	? W=184		+Ir=192.7; † Os=195;
11	-Au=197.5	-Hg=200	-Tl=204	-Pb=207	-Bi=210			+Pt=197.5
12				? Th=231		+U=240		

* The new metal, Norwegium, recently discovered by Dahll, appears to belong to this position.

† Senbert.

are always para-magnetic, whereas those elements belonging to odd series are always diamagnetic.

This is readily seen from the table above, in which the elements are arranged together with their atomic weights as in Mendeljeff's original memoir. The sign (+) signifies that the element to which it is attached is para-magnetic, the sign (-) that it is diamagnetic, and (?) that its magnetic properties in the free state are unknown.

At present I am unable to offer any explanation of these facts.

The specific magnetic powers of very few elements have as yet been determined, but judging from the few cases in which there are available data, it appears that as regards the odd elements of a group they become more and more diamagnetic as the atomic weight increases, thus:—

Specific Magnetic Powers.

Bequerel.

S = -1.14	P = -1.64	H = -0.001 (Faraday)
Se = -1.65	Sb = -4.10	Cu = -1.68 (Bequerel)
	Bi = -22.67	Ag = -2.32
		Au = -3.47

In the eighth group, also, the para-magnetism appears to diminish as the atomic weight increases, for Fe, Ni, and Co are strongly para-magnetic, whereas Rd, Pd, Pt, Os, and Ir are but feebly so. According to Faraday, also, Pd is more magnetic than Pt and Os.

In conclusion, it follows that an examination of the magnetic properties of newly discovered, and of little known, elements would render considerable service in the determination of their positions in the general classification.

The Owens College, Manchester.

ON THE NASCENT STATE OF BODIES.

By Dr. T. L. PHIPSON, F.C.S., &c.

THE note published in the last number of the CHEMICAL NEWS, by Dr. Tommasi, of Florence, demands a few words of remark.

Many years ago I showed that all bodies probably were susceptible of taking what is called the *nascent state*, a state in which their *electric* properties and powers of entering into combination are exalted to an extraordinary degree.

It would be rash to conclude from the single fact of the non reduction of perchlorate of potash quoted by Tommasi that the nascent state of hydrogen does not exist. Why should he draw such an abrupt conclusion from the investigation of some reactions given by one salt? Would it not have been as logical to conclude that nitrate of silver will not precipitate chlorine because when added to the said perchlorate it produces no precipitate of chloride of silver? Why did he not extend his researches to permanganate, &c.? I quite agree with him that the molecular notions of M. Wurtz fail to explain the phenomena in question.

In 1858 I brought forward a series of experiments in my paper on "La Force Catalytique, ou Etudes sur les Phenomenes de Contact," to which the Dutch Society of Sciences of Harlem awarded its gold medal, experiments which show that when oxygen and hydrogen combine in presence of platinum, iridium, phosphorus, humus, and other bodies, or when hydrogen precipitates gold from its perchloride in presence of a *third body*, an electric phenomena occurs in every case, and that we have here an exact reproduction of what takes place in the ordinary voltaic battery; that these substances are then in an exalted state identical with that known as the "nascent" state, that is, producing the same reactions.

A discussion arose on this subject some twelve or fourteen years ago at the Academy of Sciences at Paris, and the views put forward on that occasion by the elder Becquerel, by H. Deville, and others, showed that my interpretation of these so-called "catalytic" phenomena had been entirely adopted.

The nascent state of bodies is, therefore, no myth, as Prof. Tommasi would endeavour to make out for hydrogen. It is that peculiar exalted state which bodies assume as they enter into combination and as they leave a combination, a state in which their *physical* and *chemical* properties (especially their *electric* properties) are very different from what they are in the same bodies in a free state. Dr. Tommasi's experiments are very interesting as pointing once more to the peculiar nature of the salts formed by the oxacids of chlorine, but they throw no light on the *nascent* state of bodies.

ON A

POSSIBLE CAUSE OF VARIATION IN THE PROPORTION OF OXYGEN IN THE AIR.

By E. W. MORLEY, M.D., Ph.D.,

Professor of Chemistry in Western Reserve College, Hudson, Ohio.

PROFESSOR LOOMIS had proposed the theory that certain great and sudden depressions of temperature at the surface of the earth are caused, not by the transfer of cold air from higher to lower latitudes, but by the vertical descent of air from cold elevated parts of the atmosphere. The evidence supporting this theory was published in the *Am. Journ. Sci.* in January and July, 1875. It occurred to the writer some time since that if this theory were true, as the evidence makes very probable, the air at the surface of the earth during such a great and sudden depression of temperature might well contain a smaller proportion of oxygen than the average. Dalton reasoning from the fact that oxygen has a greater specific gravity than nitrogen, argued that the proportion of oxygen to nitrogen in the atmosphere should decrease with increasing altitude above the earth's surface; whether he clearly enough recognised that such a regular decrease would be realised only in an atmosphere in a state of equilibrium undisturbed by convection currents the writer does not know,

not having seen Dalton's memoir. Such a decrease of atmospheric oxygen with increasing altitude has not yet been detected by analysis, although the amount of decrease, on the theory that oxygen and nitrogen are distributed in the atmosphere according to the law which would prevail in case of equilibrium, is so rapid that it would be detected with ease, even in altitudes attained in every holiday ascent of a balloon. This decrease may be calculated from the formula $R = R_0 \cdot 0.9832960^H$, where H denotes the height above the earth's surface expressed in kilometres, R_0 denotes the ratio of the tension of oxygen to that of nitrogen at the surface of the earth, and R denotes the same ratio for the height H. The constant is computed from the determinations by Regnault of the weights of a litre of oxygen and of nitrogen, and of the specific gravity of mercury. The following table gives in the second column the ratio of the tension of oxygen to that of nitrogen at the height in kilometres given in the first column, and the percentage of oxygen at the same height in the third column. The percentage of oxygen at the earth's surface assumed in the table is that used in the tables for gas analysis in Bunsen's Gasometrische Methoden.

It will be seen that the composition here calculated for a height of a single kilometre is so different from that at the surface that analysis of no very refined accuracy would detect the variation with ease. But no such variation has been detected even in samples of air collected at the greatest elevations attainable.

Height, kilometres.	Ratio of O to N. Per cent.	Per cent of O.
0	26.52	20.96
1	26.08	20.68
2	25.64	20.41
3	25.21	20.14
4	24.79	19.87
5	24.38	19.60
6	23.97	19.34
7	23.57	19.07
8	23.18	18.82
9	22.79	18.56
10	22.41	18.31
20	18.93	15.92
30	16.00	13.79
40	13.52	11.91
50	11.42	10.25
60	9.65	8.80
70	8.16	7.54
80	6.89	6.45
90	5.82	5.50
100	4.92	4.69

But although this is the case, it is certain that in the atmosphere of the same place at different times the oxygen varies by more than one-fortieth of its average amount. Variations so large as this are rare, but variations of the one-hundredth or two-hundredth part are common. It therefore seemed to the writer proper to examine whether facts bear out the conjecture that certain great and sudden local depressions of temperature are caused by the descent of cold air from the upper part of the atmosphere, and that such air may by its poverty in oxygen throw some light on a question in meteorology and a question concerning the physics of a mixture of different gases.

In the number of *Wiedemann's Annalen* for April of the current year, Jolly has published the results of numerous and very accurate analyses of atmospheric air. He asserts a connection between the variations in composition detected and the direction of the winds when the sample was collected. He considers himself justified in concluding that the atmosphere of tropical regions is poorer in oxygen than that of polar regions, and supposes that at the equator more oxygen is consumed in processes of oxidation than is set free by those of reduction, while the opposite is true near the poles. Since no difference in the composition of the atmosphere at the equator and at the poles has been detected, while on this theory the difference

must be one large enough to account for the extreme variations found in temperate regions, and to account for them after such abnormal air had been exposed to admixture with air of a different composition during a passage of thousands of miles, the writer fears that the theory of Jolly will need further proof. Other reasons for a similar doubt will suggest themselves.

On the writer's theory, a sample of air collected at the centre of an area covered by a descending current of cold air would at some given instant be a sample fresh from the upper part of the atmosphere, but little exposed to admixture on the way. If before its descent it had remained at a great height for a long time, it might well have lost some of the oxygen which it contained when it was at some previous time at the level of the sea, and the difference might be enough to be detected. An observer at one side of this central point would have samples more or less mingled with surface air; but even then, a deficiency of oxygen might be detected by accurate analysis.

The writer hopes to make arrangements for the regular collection of samples at points which Professor Loomis has indicated as regions of frequent descent of cold air from great heights. But while laying plans for the work, he has thought best to ascertain whether some light on the change in the constitution of the atmosphere might be obtained by analysing samples of air collected at home. Having an apparatus for gas analysis lately constructed for the study of the gas issuing from the numerous gas wells of his vicinity, he used this for such determinations. In general, the apparatus is constructed on the plan of McLeod's modification of Frankland and Ward's apparatus. But some important modifications have been introduced, and excellent workmanship was bestowed on details. Some such points but slightly concern analysis by explosion. The connection between the eudiometer and absorption-tubes is novel, and has worked well. This will be described in some proper connection. Here will be described everything necessary for a judgment of the accuracy of the analysis to be cited.

The eudiometer and pressure tubes were made from the writer's drawings by Geissler, whose recent death is a loss to science, and a personal loss to so many who have been aided by him. The stop-stocks at the top of these tubes will retain a Torricellian vacuum for weeks. The internal diameters of the tubes are 20.9 and 10.7 millimetres. At the lower end of the eudiometer tube is a glass stop-cock, the use of which is simply to permit the ready cleaning of the tube. Its glass plug is withdrawn, and in place of it is put one of vulcanite so bored that acids or water can be aspirated through the eudiometer without dismounting it and without drawing off the mercury from the pressure tube. The stop-cock at the top of the eudiometer tube is also provided with a similar plug for the same purpose. The pressure and eudiometer tubes are surrounded by a stream of water entering at the top of the pressure tube and running away from near the bottom of the two. The level of water is kept constant by a device similar to that of Thomas, described in a late number of the *Journal of the Chemical Society*, but perhaps superior to his in some respects. The flow of mercury to and from the movable reservoir is controlled by an iron stop-cock which is attached to the iron tripod support of the whole apparatus. The plug of this stop-cock is vertical, and is prolonged by a shaft which puts it within easy reach of the observer. By means of a long handle on this shaft, the stop-cock can be moved with the greatest delicacy. From this stop-cock, an iron tube, cast in the same piece, extends under the ends of the glass tubes of the apparatus, and two small iron tubes rise from this horizontal tube; these last meet the glass tubes and are connected with them by short tubes of patent black rubber containing no free sulphur. The connectors are tied so as to endure the pressure of mercury having a head of several feet, and are surrounded with mercury so as to be absolutely air-tight. The plug of the iron stop-cock is also so surrounded with mercury that the entrance of air is ab-

solutely impossible, and the same precaution was taken at the junction of the two small iron tubes with the horizontal tube. The cast iron of this tube and stop-cock is well janned, and no leakage through its pores has occurred.

The measurement of the volume of gas in such an apparatus demands an accurate adjustment of the level of the mercury in the eudiometer tube to one of the marks of the graduations. Such an adjustment can be accurately made by admitting mercury very slowly from the reservoir, and closing the stop-cock at the required moment. But if now the temperature of the gas be not quite constant, the adjustment can be renewed for a second reading only by letting in or out a column of mercury of several millimetres, again permitting it very slowly to approach the proper level, and stopping at the instant of contact. It is quite impossible to open the stop-cock, admit the twentieth or fortieth of a millimetre of mercury, note the right instant, and then again close the stop-cock. But for accurate work, the means of altering the level of the mercury by such small quantities, and of doing it by a continuous movement, seemed important. In the end, therefore, of the horizontal iron tube, there works a plunger, packed with great care, which can be moved in or out by a screw. By means of this micrometric movement, the level of the mercury which has been adjusted by the stop-cock can be altered with the greatest delicacy, and re-adjusted till perfect steadiness is attained. Danger of leakage through the washers around the plunger was prevented by providing a seat into which, when the plunger is screwed quite home, it fits so as to cut off connection between the pressure tube and the rest of the apparatus. The mercury in the pressure tube is, by the use of this valve, always kept at such a height that any possible leakage is that of mercury outward, and not of air inward.

The eudiometer and pressure tubes are cemented into the brass cap of the glass cylinder containing the water intended to keep all parts of the apparatus at the temperature. The level of the two graduations with respect to each other is therefore constant.

(To be continued.)

NOTES OF WORK BY STUDENTS OF PRACTICAL CHEMISTRY

IN THE
LABORATORY OF THE UNIVERSITY OF
VIRGINIA.

No. VIII.

Communicated by J. W. MALLET,
Professor of General and Applied Chemistry in the University.

1. (46.) Analysis of Volcanic Ash from a Recent Eruption of Cotopaxi. By J. R. SANTOS, of Guayaquil, Ecuador.

The material for this analysis was sent to Senor Santos by a member of his family living at Bahia de Caraquez, on the coast of Ecuador, at which place it fell in large quantity on August 23, 1878. The distance from Cotopaxi is about 120 miles. The eruption of the volcano is reported as having been distinctly seen from Guayaquil.

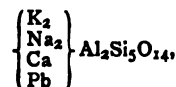
The ash was very fine, of a brownish grey colour, and appeared under the microscope to consist of nearly uniform, transparent, glassy granules, with a small admixture of reddish brown ferric oxide. It contained no particles sensibly attracted by the magnet. Heated in the blowpipe flame it fused with intumescence to a brownish yellow bead. Hydrochloric acid dissolved the oxide of iron without materially affecting the rest of the ash. Fused with borax, the reaction for iron was obtained. Lead was detectable only after decomposition by fusion with an alkali, and appears therefore to have existed as silicate. Due precaution was used to prove that it was not derived from any of the reagents or vessels employed.

Specific gravity of the ash, properly deprived of air, = 2.743. The analysis gave—

SiO ₂	..	..	..	..	..	56.661
Al ₂ O ₃	..	..	..	..	..	19.398
Fe ₂ O ₃	..	..	..	..	..	7.523
PbO	..	..	..	..	..	0.575
CaO	..	..	..	..	..	6.229
MgO	..	..	..	..	..	trace
Na ₂ O	..	..	..	..	..	6.123
K ₂ O	..	..	..	..	..	2.425
H ₂ O	..	..	..	..	..	0.862

99.796

Neglecting the ferric oxide, most of which at any rate was undoubtedly mixed with the silicate, and throwing out also the water as non-essential, the above figures approximate to those for certain lavas and trachytes, and lead pretty nearly to a formula of felspathic type,—



but it is by no means certain that there may not have been more than one silicate present.

For the sake of comparison, the following analyses of volcanic ashes, already on record, may be quoted:—

	Vesuvius (1822). Dufrénoy.	Etna (Crater).	Etna (Catania).	Hecla (1845). Genth.	Hecla (1845). Ashes on Ork- ney. Connell.	Gunung Guntur (Java). Schweizer.
SiO ₂ ..	53.67	48.737	46.309	56.89	59.20	51.64
Al ₂ O ₃ ..	17.94	17.886	16.846	14.18	15.20	21.89
Fe ₂ O ₃ ..	—	12.756	9.850	—	9.60	—
FeO ..	5.75	—	4.430	13.35	—	10.79
MnO ..	—	—	—	0.54	—	—
CaO ..	1.92	5.495	10.276	6.23	4.82	9.34
MgO ..	7.15	2.534	5.439	4.05	0.60	3.42
Na ₂ O ..	9.55	4.502	3.340	2.35	—	2.92
K ₂ O ..	4.02	2.045	1.411	2.64	6.74	0.55
Cl and SO ₃	—	—	2.207	—	—	—
H ₂ O ..	—	6.630	—	—	—	0.60
	100.00	100.585	100.108	100.23	96.16	101.15

The occurrence, in the ashes from this Cotopaxi eruption, of lead to the extent of more than 1-200th of the whole mass, which must in the aggregate have been enormously large, is interesting, as is also the different state of combination in which the metal occurs from the already known, though comparatively rare, chloride (Cotunnite) of Vesuvius, eruptions of 1822 and 1855.

2. (47.) Chemical Composition of "Livingstonite" from Huizaco, Guerrero, Mexico. By F. P. VENABLE, of Charlottesville, Virginia.

This mineral was described by Mariano Bárcena, of the City of Mexico, in 1874*, and named by him in honour of the African traveller, Dr. Livingstone. According to the original account it resembles in general aspect native antimony sulphide, with which it is apparently isomorphous. It is found in columnar groups and imperfectly detached prisms of bright lead-grey colour, with red streak, instead of the black streak of stibnite. Hardness = 2. Sp. gr. at 16° C. = 4.81. Fuses at the first touch of the blowpipe flame, giving off copious white fumes. Not sensibly attacked by cold nitric acid, but dissolved by the heated acid, leaving a white residue. Affords the reactions of sulphur, antimony, and mercury. Analysis gave Senor Bárcena—

* "Naturaleza," iii., 35 and 172. Amer. Journ. Sci., August, 1874; January, 1875, 64.

Sulphur	29.08
Antimony	53.12
Mercury	14.00
Iron	3.50
	99.70

Whence the atomic ratio was deduced, S : Sb : Hg : Fe = 18.17 : 8.7 : 1.4 : 1.2, or, nearly, 15 : 7 : 1 : 1, with the formula $4Sb_2S_3 + HgS + FeS_2$, involving an excess of 1 atom of antimony. This ratio, according to my calculation, should rather be 13 : 6.2 : 1 : 0.9, or 13 : 6 : 1 : 1. The mineral occurs at Huitzuco, in the State of Guerrero, in a matrix of carbonate and sulphate of calcium, along with native sulphur, cinnabar, valentinite, and stibnite.

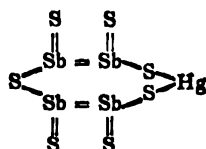
Recently Senor Bárcena has sent me some specimens of this mineral, stating that he has reason to think them purer than those formerly analysed by him, and requesting me to revise the question of chemical composition, about which he entertains some doubt. The material received agrees in all respects with the above description. The choicest portions have been carefully analysed by Mr. Venable. It was ascertained that even in these selected fragments calcium sulphate and free sulphur existed, the former soluble in water or weak hydrochloric acid, the latter in carbon disulphide. The ore was dissolved by prolonged digestion with nitro-hydrochloric acid. In one experiment, the antimony thrown down as sulphide was deprived of surplus sulphur by heating in an atmosphere of carbon dioxide : in another, sulphur was determined by oxidation with nitric acid and precipitation as barium sulphate. From one specimen the total sulphur was gotten by oxidation in the liquid way ; from another by fusion with sodium nitrate and carbonate. The quantity of mercury present was checked by distillation of a separate portion with lime. The results were—

	I.	II.
S (combined)	20.43	19.64
S (free)	3.97	3.57
Sb	46.49	44.26
Hg	19.56	18.47
Fe	0.16	0.10
$CaSO_4 \cdot 2H_2O$	7.55	12.59
Insoluble residue ..	0.52	
Moisture	0.89	
	99.57	

It is manifest that iron is not an essential ingredient. Deducting this and the calcium sulphate, free sulphur, &c., of the matrix, and re-calculating percentage, we have the following figures :—

	I.	II.	Mean.
S	23.62	23.84	23.73
Sb	53.76	53.74	53.75
Hg	22.62	22.42	22.52
	100.00	100.00	100.00

Hence the atomic ratio S : Sb : Hg = 7 : 4 : 2 : 1, which gives the formula $HgS_2Sb_2S_3$, or—



This is remarkable as representing the most strongly acid sulph-antimonite as yet known, since zinkenite contains Sb_2S_3 but once for PbS . The corresponding oxygen salt would be intermediate between the mono- and tri-antimonites of Terrell.

3. (48.) Analysis of Magnetite exhibiting unusual Crystalline Distortion, from Henry Co., Virginia. By EDWD. C. SMITH, of Richmond, Va.

The mineral occurs in loose crystals, for the most part from 15 to 30 m.m. through, on the land of Mr. J. P. Barksdale, in Henry County. There is a thin crust of ferric hydrate on the outside, but most of this is easily brushed off. The greater part of each crystal has the ordinary black colour and general appearance of magnetic iron. Hardness = a little under 6. Sp. gr. = 4.98. Strongly magnetic, but with no very distinct polarity. Many of the crystals are curiously distorted by minute step-like projections or depressions of parts of the surface, like those of some diamonds, fluor-spar cubes, &c., so that the general aspect is that of a rhombic octahedron, but with irregularly varying inclination of the general surfaces to each other, such angles as $107^\circ-15'$, $104^\circ-10'$, $100^\circ-55'$, 102° , $104^\circ-18'$, $109^\circ-30'$, $106^\circ-05'$, $102^\circ-48'$, and 115° , being obtainable with the common goniometer of application. So minute are the successive steps that the appearance is strongly suggestive of true crystalline form other than monometric, and pseudomorphism was at first suspected. A certain amount of irregular pitting of the surface is also observable under the crust, the result of chemical change to ferric hydrate.

Analysis by Mr. Smith of the clean interior portions of the crystals gave—

FeO	30.56
Fe_2O_3	67.94
Insoluble residue (quartz) ..	1.56
	100.06

corresponding accurately to simple magnetite. There was no titanium present.

4. (49.) Analysis of Crude Rock-salt from Saltville, Washington Co., Virginia. By B. E. SLOAN, of Charleston, South Carolina.

Large supplies of salt are now, and for a long time have been, obtained in this neighbourhood, in the extreme south-western portion of the State, by boiling down brine, originally found issuing naturally from the ground, but for which exit on a larger scale has been artificially produced by digging and boring. In the course of these excavations the underlying solid salt has been occasionally exposed, though never worked on an industrial scale. A year or two ago I received from the superintendent of the works some specimens of dark brownish red crystalline rock-salt, thus taken out during the deepening of one of the salt wells, and a portion of this material has, as an example for practice, been analysed by Mr. Sloan. He found—

NaCl	89.21
KCl	trace
$CaSO_4 \cdot 2H_2O$	4.86
Fe_2O_3	0.84
SiO_2	4.53
	99.44

No strontium, barium, or lithium could be detected. Some thin translucent plates of ferric oxide were visible under the microscope in the residue insoluble in water, but nothing like the beautifully regular crystalline spangles obtainable from the Carnallite of Stassfurt.

5. (50.) Analysis of "Tungsten-Manganese Bronze." By F. P. VENABLE, of Charlottesville, Va.

Amongst a number of preparations of tungsten lately received from E. W. L. Biermann, of Hannover, were sundry alloys purporting to contain this metal in union with others. As two or three cases have of late years

been published of alloys supposed to contain the more refractory metals, as tungsten, chromium, titanium, &c., and for which special merit was on this account claimed, while little or none of the metals in question could be found on analysis, it seemed worth while to examine at least one or two of these German specimens. A fragment cut from a small block of so-called tungsten-manganese bronze, of light gold-yellow colour, pretty close grain, and with a fine polish upon one side of the specimen (sp. gr. = 8.64), was carefully tested qualitatively, and the proportions of the constituents found were thus determined. No manganese whatever was present, and but an insignificant amount of tungsten. The alloy consisted of—

Cu	86.51
Sn	9.04
Zn	3.47
Fe	0.26
W	0.23

99.51

It is therefore but an ordinary gun-metal, with part of the tin replaced by zinc. Of course manganese may have been added in its production, as well as a larger proportion of tungsten; but if so, they have failed to be taken up, or have been burnt out before the alloy was cast; and manganese so removed may have served in a measure to improve the compactness and homogeneity of the mass by carrying off with it oxygen from the remaining metals, as in the use of phosphorus in making "phosphor-bronze. But the name under which the alloy is sold is calculated to mislead purchasers.

(To be continued.)

Presence of Alcohol in Animal Tissues during Life and after Death in Cases of Putrefaction from a Physiological and Toxicological Point of View.—J. Béchamp.—MM. Schrader and Dusch have shown that well-cooked meat may be preserved for several weeks without alteration in presence of filtered air; and that meat heated in the water-bath, so as merely to coagulate the surface, became putrid under the same conditions. In order to find out the cause of this decomposition, the author performed the following experiment:—A piece of horse flesh, weighing 3 kilos., was plunged for ten minutes in boiling water, so as to coagulate the surface, and was then placed in a crystallising pan covered over with a very compact linen cloth. After the lapse of thirty-eight days the vessel was opened. A small quantity of liquid had collected, which swarmed with *Vibrio*nes. The odour of the meat was offensive, but distinct from that of ordinary putrefaction. On cutting the meat open to examine the central part the muscular striation had disappeared. Under the microscope were found a very few free *Microzymas* in the free state, and a larger number associated; a great quantity of various *Bacteria* (*B. termo*, *articulatum*, and *capitum*), and even a few *Leptothrix*, but not a single *Vibrio*, a fact proving that the air had not penetrated into the interior. From the mass the author isolated about 0.8 grm. of alcohol, the identity of which was demonstrated by the usual reactions, along with 10 grms. of sodium acetate and butyrate, and salts of acids higher than the butyric. Hence putrefaction completely approaches in its nature to fermentation properly so-called. But during life the transformations which are effected within animal tissues are due also to microzymes, and approximate consequently to fermentation. Hence alcohol should be present in the organs. Experiments on the liver and the brains of sheep and the brains of oxen clearly showed its presence. These facts have a deep interest, both physiologically and toxicologically. The presence of alcohol in the tissues can no longer be considered a proof of its ingestion previous to death.—*Comptes Rendus*, lxxxix., No. 12.

BURETTE FOR COLLECTING, MEASURING, AND DELIVERING GASES OVER MERCURY.*

By PHILIP BRAHAM, F.C.S.

THE burette consists of a divided glass tube open at both ends, the lower end, A, having a lip, the upper end being covered with a loosely fitting boxwood cap. A piston, B, formed of a disk of india-rubber between two plates of steel connected with a steel rod, C, is fitted into the tube.



To fill the burette, the piston is depressed below the lower end until a portion of the mercury escapes above it. On drawing it up any portion of the tube can be filled. When the gas has been delivered the measurement may easily be made by lowering the piston till the mercury inside is level with that on the outside, and the contents read off. By further depression of the piston, and placing the lip under a eudiometer, any required amount of gas may be delivered for analysis, the clamp E being used to stop the piston when the required amount is discharged. The clamp D is used to prevent the piston descending when the instrument is full of mercury.

* Read before the British Association for the Advancement of Science Section B., Sheffield, 1879.

NOTICES OF BOOKS.

Chemical Denudation in Relation to Geological Time. By T. MELLARD READE, F.G.S. London: David Bogue.

IN view of the recent attempts made by certain physicists to fix a maximum limit for the past existence of the earth—attempts characterised not more by their extreme assumptions than by the conflicting character of their results—geologists and biologists are naturally led to revise the evidences in favour of a far higher terrestrial antiquity. Mr. Reade, in the work before us, takes what we believe to be novel ground. By "chemical denudation" he understands the quantity of dissolved matter carried down from the land to the sea by rivers, and hence primarily by the rainfall. The proportions of carbonates and sulphates of lime and magnesia, of chloride of sodium, besides certain minor constituents, are taken from the analyses of Prof. Frankland in the "Reports of the Pollution of Rivers' Commission," and from the researches of Prof. Bischoff on the principal European rivers. The quantity of water run off by the rivers—a much smaller amount, of course, than the annual rainfall—is next taken into consideration. The remaining elements in the calculation are the total contents of the ocean, 2,494,500 billions of tons, according to Herschell, and the analysis of sea-water as given by Prof. Frankland, which shows 48.9 tons carbonate of lime and magnesia, and 1017 tons of the sulphates of the same bases; and, lastly, 3259 tons of common salt per 100,000 tons of water. Hence it would take 480,000 years to introduce into the ocean its present proportion of the carbonates of lime and magnesia, 25 million years to supply the sulphates, but 200 million years to furnish the salt. It may be incidentally mentioned that the sodium chloride existing in solution in the ocean is sufficient, if solidified, to cover the whole of the dry land with a bed of salt 914.9 feet in thickness. These 200 million years the author considers as an inferior limit.

In the second section of the work, a paper on the "Geological Significance of the Challenger Discoveries," the author points to the coldness of the bottom of the ocean at great depths as a proof of the extreme slowness of the secular cooling of the earth. He considers that the accumulation of one foot of chalky clay over the bottom of the ocean must require 20,000 years, and a foot of red clay probably ten times as long. He concludes also that even the deep seas have at one time been continents. Thus "the molluscan fauna of the eastern coast of North America is very similar to that of Europe," which could not have happened without littoral contiguity.

The remaining portion of the work is a paper read before the Royal Society on "Limestone as an Index of Geological Time." The author's results may be thus summarised:—Taking the average rainfall run off the igneous ground during all geological time at 28 inches per annum, and the amount of carbonate and sulphate of lime it takes up from the igneous rocks at 4.00 per 100,000, the annual yield of the land in those forms of lime would be 71.68 tons yearly per square mile. Calling it 70 tons, one-tenth of the area of the land, or 5,100,000 square miles, would yield 357,000,000 tons yearly of these minerals. The total area of the globe is 196,900,278 square miles, so that the land would yield to the whole globe 1.813 tons per square mile yearly. At 13½ cubic feet to the ton it would take 1,139,032 years for the accumulation of a deposit one foot in thickness. Therefore 528 feet thick, the supposed minimum thickness of limestone in the sedimentary crust of the earth, would be eliminated from the original material of the earth in 601,408,896 years.

Prof. Ramsay, from "considerations of a mixed physical and palæontological character," assigns to the groups from the Laurentian down to the Post Pliocene a total time of 600 million years.

Space will not permit us to enter into the arguments which may be presented on both sides of this important

question. But we must admit that the author has very fairly and ably met the majority of the objections which may be urged against his views.

CORRESPONDENCE.

RUSTLESS IRON.—BARFF'S PROCESS.

To the Editor of the Chemical News.

SIR,—The great interest which has been taken in Prof. Barff's process for the prevention of corrosion on iron surfaces by all persons connected with the iron trade leads us to believe that it would be interesting for your readers to know that we are now supplying all kinds of "Anti-Corrodo" iron goods by license under Prof. Barff's patent, and which obtained two Medals at the Paris Exhibition. The process, shortly, consists of passing superheated steam over the iron goods to be treated whilst at a red-heat, and can be applied to all kinds of iron-work, rendering it absolutely rustless at a less cost than galvanising, so substituting an absolute protection for one which confessedly is but partial in its action and easily removed. Amongst "Anti-Corrodo Goods," wrought-iron tubes stand out prominently, as they and a host of similar goods are peculiarly well adapted for the application of Prof. Barff's process, and we have an iron chamber 12 feet long specially built for treating this class of goods, whilst a chamber 7 ft. × 3 ft. × 13 ft., which we built to coat a large quantity of large iron railings for the Duke of Norfolk, enables us to treat all kinds of iron-work of any ordinary dimensions for manufacturers and others interested in the process.

A report of Sir Joseph Whitworth goes to prove that the iron is slightly improved in strength by the process. Many of the articles first treated are in our possession, and although they have been exposed to the severest of tests by various firms and gentlemen in whose possession they have been, in some cases over two years, they present no appearance of corrosion, and are, as claimed for them, rustless and incorrodable.—We are, &c.,

THE RUSTLESS AND GENERAL IRON CO.,
JAMES E. SPENCER.

97, Cannon Street, E.C.

PYRITES-SMALLS IN THE MANUFACTURE OF SULPHURIC ACID.

To the Editor of the Chemical News.

SIR,—A person under the name of "Sulphur" enquires as to the cheapest way of using up pyrites-small in manufacturing sulphuric acid. I beg to refer him to the excellent manual of Professor Lunge, of the Zürich Polytechnikum, recently published by Van Voorst, "On the Alkali Manufacture." He will there find an account of a simple kind of furnace or burner where the smalls are arranged on shelves, placed alternately under one another so that the ore may be gradually worked downwards, and falls at length to the lower part of the oven very satisfactorily burnt. This form of burner is used in a vitriol works on the side of the Lake of Zürich, and I know it is a success, for smalls alone are used there, to the best of my knowledge. I have myself seen this form of burner in operation, and to me it seemed to answer perfectly. Speaking from memory, I believe I was told the ore could thus be burnt down to 5 per cent of residual sulphur as a maximum.—I am, &c.,

WATSON SMITH, F.C.S., F.I.C.

Stretford, near Manchester.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 12, September 22, 1879.

Evolution in Medicine.—C. Sedillot.—Remarks on medical progress, in which the author quotes the aphorism of Hippocrates, "everything is natural and everything is divine."

Influence of Atmospheric Electricity on the Growth, Inflorescence, and Fructification of Plants.—M. C. Naudin.—The author, referring to the experiments of MM. Grandeau and Leclerc, at Nancy, in which plants of tobacco and maize were withdrawn from the influence of atmospheric electricity by being enclosed in wire cages or placed near large trees, and were found in consequence to be retarded and impoverished in their growth and fructification, describes similar experiments, which he has conducted at Antibes upon kidney beans, lettuce, tomatoes, and the herbaceous cotton-plant. The results were, however, exactly the reverse, the plants being benefitted, though to an unequal extent, by being withdrawn from atmospheric electricity. The weight of fruit obtained from the tomato was slightly more than double the weight obtained from a plant placed otherwise under conditions exactly similar. He concludes that the action of electricity, like that of heat, light, and moisture, varies upon different species of plants.

Theoretical Essay upon the Law of Dulong and Petit: Case of Perfect Gases.—H. Willotte.—A mathematical paper, not suitable for abstraction.

No. 13, September 29, 1879.

Researches on the Effects and the Mode of Action of Substances used in Antiseptic Dressings.—MM. Gosselin and A. Bergeron.—Experiments on the behaviour of blood mixed with known quantities of phenol, of alcohol at 86 per cent, and of camphorette spirit.

Theoretic Essay on the Law of Dulong and Petit.—H. Willotte.—A continuation of the memoir inserted in No. 12. The author treats here of solids, liquids, vapours, and compound bodies.

Vibratory Forms of the Bubbles of Glyceric Liquid.—C. Decharme.—The author proposes three laws:—For the same number of nodals (the same system) the diameters of the bubbles are proportional to the lengths of the vibrating plate, or inversely proportional to the square roots of the numbers of vibrations. For the same diameter of bubbles the numbers of nodals are inversely proportional to the lengths of the vibrating plates, and directly proportional to the square roots of the number of vibrations. For the same length of vibrating rods the numbers of nodals are proportional to the diameters of the bubbles.

Action of Metallic Nitrates upon Monohydrated Nitric Acid.—A. Ditte.—Metallic nitrates may act upon monohydrated nitric acid in three very different manners. Ammonium nitrate combines with nitric acid to form an acid salt, which, at ordinary temperatures, dissolves a large quantity of melted ammonium nitrate, forming another acid salt. Potassium nitrate yields an acid salt in the same manner, as do also thallium and rubidium nitrates.

Thermic Researches on Succinic Acid and its Derivatives.—P. Chroustchoff.—Succinic acid dried in the water-bath and sifted gives, on solution in 500 c.c. at 11°, an absorption of heat:— $C_8H_4O_8H_2$ —6.4 cal.

A New Curare, Extracted from a single Plant (*Strychnos triplinervia*).—MM. Couty and de Lacerda.

—The authors have obtained from the bark of this *Strychnos*—a species common in the province of Rio Janeiro—an extract, which, though weaker than that prepared by the Aborigines, gave, when introduced into the blood or under the skins of animals, the well-known symptoms of curare.

Moniteur Scientifique, Quesneville.
September, 1879.

A great part of this issue is devoted to a review of the *Splendeurs de la Foi*, of M. l'Abbé Moigno. Save the usual reports of the meetings of the Academy of Sciences, the rest of the number is taken up with Mr. Perkin's history of alizarin, and Dr. O. H. Witt's paper on "A new Class of Aniline Colours," both from well-known English sources, and with a paper on the "Reynier" electric lamp, which cannot be reproduced without the accompanying illustrations.

October, 1879.

Aniline-Black.—R. Kayser.—The author has prepared aniline-black by different methods, and has found that when pure aniline was used the base formed possessed the composition $C_{12}H_{10}O_2$, Reineck's nigranilin. This base is insoluble in water, alcohol, ether, and chloroform; soluble in creosote and aniline with a blue colouration, rapidly turning brown under the influence of heat. In sulphuric acid it dissolves with a violet colour. The sulphate and hydrochlorate are decomposable by water. The formation of nigranilin takes place in acid solutions only; the presence of a metallic salt is not necessary, especially if there is an excess of a mineral acid. The author's experiments show that toluydin does not take part in the formation of aniline-black, and that in practice a pure aniline should be used. With reference to the view of M. Nietzki that aniline-black contains two isomeric bases, the one easily soluble in chloroform and aniline with a blue-violet colour, and the other, less readily, with a brown, the author has not observed a base soluble in chloroform.—*Verein. Beförder. Gewerbfl. Wiss.*

Aniline-Black.—R. Nietzki.—The author refutes the assertion of M. Dechend that aniline-black is an oxygenated base, $C_{24}H_{17}N_3O_2$. Aniline-black, which possibly approaches the indulins, is of a violet colour when free, and green when existing as an acid salt, but these shades are sufficiently intense to pass for black.—*Verein. Beförderung. Gewerbfl. Wiss.*

The remainder of the papers in this issue have either been already noticed, or are derived from English sources.

Justus Liebig's Annalen der Chemie,
Band 198, Heft 1 and 2.

Communication from the Chemical Laboratory of Greifswald.—This consists of a memoir by Dr. O. Zander on the amido-disulpho-benzolic acids, which the author obtains by heating the three amido-sulpho-benzolic acids with fuming sulphuric acid.

Influence of Volume and Temperature in the Preparation of Ozone, and Description of a New Ozonisorator.—A. R. Leeds.—At the temperature of +6° no ozone appears to be formed. From this point the yield increases till the temperature reaches +24°, after which it again decreases in a corresponding manner. In preparing ozone by means of the slow combustion of phosphorus, the author uses, instead of water, a solution of potassium dichromate mixed with sulphuric acid, and aspires or forces a current of air over the phosphorus, as first proposed by A. Riche. For the requisite tubes he finds a certain quality of "kerite," manufactured by Mr. A. G. Day, the best material.

Communications from the Laboratory of the University of Halle.—These consist of memoirs by W. Heintz on the oxidation-products of di- and tri-aceton

amin, especially on amido-dimethyl-acetic acid, amido-dimethyl-propionic acid, and imido-dimethyl-acetic-dimethyl-propionic acid; on the chromates of tri-aceton-amin, and on the compound of platinum chloride with the hydrochlorate of urea.

Preparation of Perbromic Acid.—Dr. G. Wolfram.—The author finds that perbromic acid is not formed by Kämmerer's process.

Researches from the Laboratory of Prof. T. Zincke.—This collection comprises memoirs by M. Rhalis on ortho-brom-benzoic acid; by T. Zincke, on compounds of the hydrobenzoin and stilben series; by A. Breuer and T. Zincke, in continuation of the last paper, and on the bodies formed by the action of dilute sulphuric acid on hydro- and isohydro-benzoin; and by T. Zincke, on the isomerism between hydro- and iso-hydro-benzoin. In this memoir the author discusses the question of physical isomerism, not to be explained by a different grouping or position of molecules. He suggests the assumption of different physical molecules, consisting of identical chemical molecules.

Contribution to the Knowledge of Glyoxylic Acid.—Dr. Carl Böttinger.—The transition of glyoxylic acid into oxalic acid and glycolic acid is connected with the decomposition of a salt of definite composition. With hydrocyanic and hydrosulphuric acids, glyoxylic acid behaves similarly to pyruvic acid. By ammonia it is converted into amido-glyoxylic acid, and by aniline into anil-glyoxylic acid. Glyoxylic acid behaves as an aldehyd and may be distinguished from pyruvic acid by its less tendency to condensations, due to the absence of a hydro-carbon residue.

Carbo-hydrates of the Tubers of the Jerusalem Artichoke (*Helianthus tuberosus*).—Dr. E. Dieck and B. Tollens.—In the tubers examined little or no inulin was found, but considerable quantities of levulin and of a dextrorotatory sugar. The composition of levulin is $C_6H_{10}O_5$, as is also ascribed to gum, starch, and dextrin. It is optically inert, has a great similarity to gum and dextrin, and in contact with yeast passes into the alcoholic fermentation. If boiled with sulphuric acid it yields levulinic acid. The sugar formed from levulin reduces Fehling's solution strongly, and has a specific rotatory power to the left (α) $4D$ at $20^\circ C.$ = 52° referred to levulin, and 47° referred to sugar. In the fermented juice mannite and glycerin have been detected, and on one occasion succinic acid.

Isomeric Nitro-salicylic Acids.—H. Schiff and F. Masino.—Not suitable for useful abstraction.

On Di-iod-nitro-phenols.—R. Piria.—The author has obtained di-ido-nitro-phenol in two isomeric modifications, the one forming pale yellow rhombic prisms, and the other gold-coloured micaceous plates, like iodide of lead.

Chemiker Zeitung.
No. 37, 1879.

Alizarin Blue.—G. Auerbach.—Tissues dyed with this colour are much affected by light. An exposure to the sun for two or three hours is sufficient to convert the original bright blue shade to a dull grey. This want of permanence, joined to the high price of the colour, have greatly limited its consumption, and since the introduction of methylen blue, which dyes cotton without the intervention of a mordant, alizarin blue has disappeared from commerce. It is prepared by heating 1 part of mono-nitro-alizarin, as dry as possible, with $1\frac{1}{2}$ part glycerin, and 5 parts sulphuric acid. After the reaction is over the mass is boiled in water, yielding a deep brownish red solution, from which, on cooling, the colour is deposited as a sulphate in brown crystals. This compound, however, can exist only in a strongly acid solution. On filtration and washing with water, the brown mass turns blue, and the colour is thus obtained in the free state. It may be re-crystallised

from amylic alcohol, from glacial acetic acid, or from naphtha. The composition of alizarin blue is



If heated to 180° with acetic anhydride it yields di-acetyl-alizarin blue, an unstable compound.

No. 38, 1879.

In connection with the general Congress of German Apothecaries, held at Hanover on September 4, was exhibited a collection of pharmaceutical antiquities, including a specimen of genuine oriental bezoar, a goblet of metallic antimony, formerly used for imparting a purgative effect to wine or beer, a venerable sample of "Album Græcum," &c., a selection of chemical apparatus inscribed with alchymistic symbols, a gaily-adorned jar for containing "Mithridate," &c.

Atti della R. Accademia dei Lincei.
Fascicolo 7, 1879.

Theorems on the Distribution of Constant Electric Currents.—Prof. G. Ferraris.—Briot, in his *Traité de la Chaleur*, having observed that in filiform conductors traversed by constant galvanic currents, as the law of Joule is a consequence of the law of Ohm, reciprocally the latter is a consequence of that of Joule, Ferraris generalises this theorem for conductors of any form soever.

Crystallographic Studies on Certain Bodies of the Aromatic Series.—Giuseppe la Valle.—Not capable of useful abstraction.

Glycerin and the Pancreatic Digestion.—Prof. A. Herzen.—The author, in opposition to Prof. F. Lussana, shows that the pancreatic liquid is pre-eminently the digestive secretion, acting upon the albuminoids as well as upon amylaceous and fatty bodies. He also refutes the objection that the albumin is dissolved by glycerin used for the extraction of the pancreatic secretion.

Nature of the Specific Agent which Produces Malarial Fever.—Prof. Edwin Klebs.—The cause of malaria is not equally developed in soils of like composition and equally moist, and it may hence be suspected to be a specific organism, requiring for its development not merely favourable outward circumstances, but also the presence of a germ. From the air of malarial districts aspirated through suitable apparatus the author has obtained organisms of about 0.95 micro-millimetres in maximum diameter, capable of producing malarial fever on inoculation. For these the author proposes the name *Bacillus malaria*. They are developed in presence of free oxygen, and belong therefore to the *Aerobii* of Pasteur.

Thermic and Galvanometric Laws of the Electric Sparks Produced on the Complete, Incomplete, and Partial Discharge of Condensers.—Prof. Emilio Villari.—The following laws are laid down:—The galvanometric deviations produced by incomplete discharges are proportional to the quantity of electricity which produces it. The heat caused by the sparks generated on an incomplete discharge is proportional to the quantity of electricity which produces it.

Vapours Diffused within Liquids.—S. Cantoni.—An extensive and mathematical paper, not admitting of useful abridgment.

Corundiferous Felspar of Biellese.—S. Sella.—The composition of this mineral is:—

Alumina	93.725
Ferric oxide	1.094
Silica, with titanic acid	3.141
Lime	traces
Water	0.867

98.827

Singular Decomposition of Hydrochlorate of Phenyl-ethyl-amin.—M. Fileti and A. Piccini.—Noticed under *Gazzetta Chimica Italiana*.

On Digallic Acid and on the Alleged Artificial Tannin.—Dr. Freda.—Also noticed under *Gazzetta Chimica Italiana*.

Certain Derivatives of Santonin.—G. Carnelutti and S. Cannizzaro.—By boiling santonin with hydriodic acid and phosphorus a powerful monobasic acid is produced, to which the authors give the name of the santonosic.

A New Isomer of Santonin.—S. Cannizzaro and L. Valenti.—The compound, on treatment with a boiling solution of caustic potassa, is transformed into a meta-santonate.

Chemico-microscopic Researches on Ashes from Etna, falling at Reggio in Calabria, May 28, and on Lava collected at Giarre, June 2.—S. Cossa.—Not suitable for abstraction.

Temperature of the Voltaic Arc and its two Polar Extremes.—S. Rosetti.—The temperature of the positive pole is greater than that of the negative. The extreme point of the former has a temperature of about 4000°, whilst the corresponding negative point scarcely exceeds 3000°.

Gazzetta Chimica Italiana.
Anno ix., 1879. Fasc. 6 and 7.

Tungsten Chlorides and Oxychlorides.—Ugo Schiff.—By the action of phosphorus chloride upon tungstic anhydride there are formed in the first reaction oxychlorides, and especially oxy-tetra-chloride. These oxychlorides to an elevated temperature or pressure, and to the action of an excess of phosphorus chloride, pass into hexachloride, which is then partially decomposed into chloride and penta-chloride.

Process for the Economical Preparation of Bibasic Quinine Citrate.—F. Dotto-Scribani.—The author dissolves in three litres of boiling water acidified with 3.669 grms. monohydrated sulphuric acid, 100 grms. bibasic quinine sulphate, and adds to the solution 32.685 grms. of tribasic calcium citrate.

Researches on Satureia Juliana.—Dr. P. Spica.—This plant is used by the country people in some parts of Italy as a remedy for intermittents. The author has extracted from it a non-nitrogenous principle, to which he assigns the formula $C_{35}H_{56}O_4$.

Chemical Study on the Salts from the Mother-liquors of the Saline Springs of Volterra.—Dr. A. Funaro.—The author has examined these waters as a commercial source of the salts of potash.

Analysis of a Chrysocolla from Chili.—Niccolo Pellegrini.—In different portions of the ore the oxide of copper ranges from 31.913 to 65.306 per cent.

Singular Decomposition of the Hydrochlorate of Phenyl-ethyl-amin.—M. Fileti and A. Piccini.—The hydrochlorate of phenyl-ethyl-amin if heated in a small tube sealed at the lamp undergoes a decomposition; on opening the tube there is not the slightest escape of gas, but there is perceived a slight odour of an aromatic hydrocarbon, and there is formed a hydrochlorate less soluble in water than the original substance. If the temperature is raised to 270° to 280° the quantity of the sparingly soluble hydrochlorate decreases, but the aromatic odour is stronger, and on adding water oily drops are separated.

Certain Neutral Ammoniacal Salts.—Fausto Sestini.—The salts in question are the citrate, phosphate, and photo-santonate.

New Experiments on Resins.—Giacomo Luigi Ciamician.—Abietinic acid and colophonium distilled with zinc powder yield toluol, meta-ethyl-toluol, naphthalin, methyl-naphthalin, and methyl-anthracen. Resin of elemi distilled in the same manner produces toluol, meta- and para-methyl-ethyl-benzin, and ethyl-naphthalin. Gum ammoniacum similarly treated gives very different pro-

ducts, one of the principal being $C_{13}H_{20}$, a hydrocarbon of the benzol series. It is probable that all the resins which resist the action of fused caustic potash are decomposed on reduction with zinc powder into hydrocarbons belonging to the naphthalin series.

Isomeric Nitro-salicylic Acids.—Ugo Schiff and F. Massino.—Not adapted for abstraction.

Alleged Artificial Tannic Acid.—Dr. Pasquale Freda.—By the action of arsenic acid, gallic acid, whether in alcoholic or aqueous solution, is not transformed into digallic acid, but into an arsenical compound having some of the properties of tannic acid. If this compound is freed from arsenic by means of hydrogen sulphide the gallic acid reappears unaltered.

On Piperidin.—R. Schiff.—A short notice of some bromine derivatives of this base.

Action of Potassium Cyanide upon the Ammoniacal Derivatives of Chloral.—R. Schiff and S. Speciale.—The compounds thus submitted to the action of potassium cyanide are chloral-ammonium, chloral-acetamid, and chloral-benzamid.

Crystalline Form of Anglesite from Sardinia.—Quintino Sella.—Upwards of seven pages of figures and crystallographic formulæ.

Crystalline Form of Certain Substances of the Aromatic Series.—Dr. R. Panebianco.—Incapable of abstraction.

Lithofellic Acid and Certain Lithofellates.—Dr. G. Roster.—An account of the preparation of lithofellic acid and of its sodium and barium salts, with particular notice of their crystalline forms.

Researches on the Lavas of the Volcanoes of Ernici in the Valley of Sacco.—S. Speciale.—The lavas contain alumina, magnesia, lime, potash, soda, ferrous and ferric oxides, traces of copper, silica, phosphoric acid, and traces of carbonic acid with water, besides imponderable traces of manganese, and of barium and lithium, visible with the spectroscopie.

Detection of Nitric Acid in Presence of Nitrous Acid.—A. Piccini.—The substance is dissolved in water along with a good quantity of urea, and the solution is added little by little to another solution of urea in dilute sulphuric acid. When the decomposition is complete some ioduretted starch paste is added. If the colourless liquid takes a blue colouration on the addition of a fragment of zinc, this indicates the presence of nitrates.

La Correspondance Scientifique.

No 61, September 23, 1879.

Sophistications of Olive Oil.—The adulteration of this oil has become so prevalent that the Minister of Agriculture and Commerce has requested the Academy of Sciences to ascertain the most trustworthy method for the detection of such frauds. Among the procedures at present under examination by a special committee is the use of the diagometer, an instrument devised by Prof. Luigi Palmieri, founded on the difference of the electric conductivity of oils. Seed oils are as a class better conductors than olive oil. At the same time every oil conducts the better the greater are its impurities. Linseed and cotton seed oil are among the best conductors, whilst the oils of pine seeds and of hazel nuts are almost as feebly conductive as the purest olive oil, known in commerce as virgin oil. Fortunately these two oils are too rare and costly to be used in the adulteration of olive oil. The use of the diagometer requires considerable manipulative skill.

Reimann's Färber Zeitung,

No. 35, 1879.

At a recent meeting of the Berlin Dyer's Association the new colour, "Puteaux blue," manufactured by MM.

Patry, of Puteaux, came under discussion. It was considered to be in all probability an indulin, but was pronounced not well adapted for wool-dyeing. For mixed colours upon silks, and as a ground for blues it was said to be applicable. Dr. Reimann, in speaking of "pittacall blue," considered that at no very distant period the products of beech tar would be utilised for the production of artificial tannin.

The "Weighting" of Silks.—Marius Moyret.—The system of weighting is ruinous to the silk-growing departments of France. Their high-class products are no longer in demand, as inferior and cheaper foreign silks serve equally well for loaded tissues. Hence these districts, already suffering from the phylloxera and from the loss of the madder trade are in the utmost distress. The author proposes that in the sale of silks, as in that of gold and silver, the proportions of the real article and of weighting matters should be exactly specified, and that a central office for the cheap and rapid assay of silks should be opened in Lyon. He states that the most excessive weighting has been carried out by a New York firm.

MISCELLANEOUS.

Lightning Conductors.—The delegates nominated in 1878 by the Royal Institute of British Architects, the Society of Telegraph Engineers, the Physical Society, and the Meteorological Society, "To consider the possibility of formulating the existing knowledge on the subject of the protection of property from damage by electricity, and the advisability of preparing and issuing a general code of rules for the erection of Lightning Conductors," have already collected a large amount of thoroughly practical information. Being anxious, however, that their Report should be as trustworthy and as exhaustive as possible, they invite correspondence upon the following points:—Full details of accidents by lightning, stating especially whether the building struck had a conductor or not. If there was a conductor, state its dimensions, construction, mode of attachment to the building, whether its top was pointed, distance of its upper terminal from the place struck, nature and extent of the connection between the conductor and the earth, and whether the earth was dry or moist, whether the conductor was itself injured, and whether the conductor or the point struck was the most salient object in the vicinity. Information is also desired, either verbally or by sketches, as to the position of metal spouting and lead roofing relatively to the point struck, and to the conductor. Details of the thickest piece of metal melted by a flash of lightning are much needed. Unimpeachable evidence of the failure of conductors is much desired, as such failures would be extremely instructive. The Secretary to the Conference is Mr. G. J. Symons, F.R.S., 30, Great George Street, S.W.

Sanitary Institute of Great Britain.—The Proceedings of the Autumn Congress of the Sanitary Institute, which will be held at Croydon, will be as follow:—Tuesday, October 21, 1 p.m., Public Luncheon; 3 p.m., Opening of Exhibition; 8 p.m., First General Meeting, Opening Address, by Benjamin W. Richardson, M.D., F.R.S., President of the Congress. Wednesday, October 22, Section I., "Sanitary Science and Preventive Medicine;" Address by Alfred Carpenter, M.D., President of the Section; Papers and Discussions, on "Sanitary Science and Preventive Medicine;" 8 p.m., *Conversazione*. Thursday, October 23, Section II., "Engineering and Sanitary Construction;" Address by Captain Douglas Galton, R.E., C.B., F.R.S., President of the Section; Papers and Discussions, on "Engineering and Sanitary Construction;" 8 p.m., Lecture to the Congress, by Prof. W. H. Corfield, M.D. Friday, October 24, Section III., "Meteorology and Geology;" Address by G. J. Symons,

F.R.S., President of the Section; Papers and Discussions, "Meteorological, Geological, and Geographical;" 5 p.m., Closing General Meeting of Congress; 7 p.m., Public Dinner. Saturday, October 25, Excursions. The Lectures to the Congress and the General Meetings will be held in the Large Public Hall, and the Sectional Meetings in the Small Hall adjoining. The Exhibition will be held in a Marquee erected at the Central Croydon Station, and will remain open until Saturday evening, November 8.

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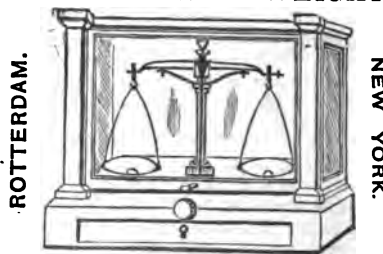
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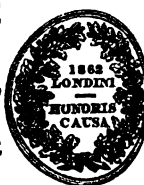
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THE CHEMICAL NEWS.

Vol. XL. No. 1039.

AGRICULTURAL CHEMISTRY IN JAPAN.

By ROBERT WARINGTON.

In the International Exhibition, now being held at Sydney, a collection of soils, manures, and agricultural products is shown by the Imperial College of Agriculture in Tokio, Japan. Accompanying the collection is a descriptive catalogue, compiled by Mr. Kinch, the Professor of Chemistry at the College, in which a short account is given of the various products exhibited, with chemical analyses of the majority of them. The catalogue contains in all about 80 analyses of Japanese products. The whole of this important work has been accomplished in the laboratory of the College, under the superintendence of Mr. Kinch, and is certainly highly creditable to the Institution. We propose to select from this Japanese chapter in agricultural chemistry some of the results which appear to have a special interest.

The catalogue opens with some analyses of soil. Then follow analyses of manures, including lime, wood ashes, nitre, waste vegetable substances, and residues from various manufactures, fish manure, bone, superphosphate, bird's dung, and hair. Next in order come analyses of foods.

Rice.—This is the grain principally cultivated in Japan. The composition of ordinary rice, and of the glutinous rice used in preparing the New Year's cakes, is as follows:—

	Ordinary Rice.	Glutinous Rice.
Water	13.63	12.01
*Albuminoids	5.80	5.13
Fat	2.15	3.30
Starch, &c.	73.14	73.20
Fibre	4.00	4.91
Ash	1.28	1.45

Mr. Kinch remarks that other specimens of Japanese rice gave 11.0 to 14.0 per cent of water, 6.07 to 7.77 per cent of albuminoids, and 1.08 to 1.33 per cent of ash.

Soy Bean (*Glycine [Soja] hispida*).—This bean is remarkable for containing a very high percentage both of albuminoids and fat: it forms probably the most concentrated food furnished by the vegetable kingdom.

	White Soy Bean.
Water	11.32
Albuminoids	37.75
Fat	20.89
Starch and soluble cellulose	24.58
Fibre	1.50
Ash	3.86

A variety of foods and condiments are prepared from this bean. By mixing the boiled beans with fermenting rice and some salt, and allowing the whole to stand in a cool place for a month, the food known as *Miso* is prepared. By precipitating the watery extract of the boiled beans with common salt, a crude legumin rich in fat is manufactured; this is known as *Tofu*.

Shoyu, known in this country as Soy, is made by a complicated process from soy beans and wheat. These ingredients are roasted, then mixed with fermenting wheat, and kept for some days in a warm room till covered with a fungus. The mass is then extracted with a solution of common salt, and the mashing preserved in vats for three

or five years. It is then filtered for use, honey or fermented rice being sometimes added if a sweet soy is desired.

Sweet Potato (*Batatas edulis*).—The tubers were found to have the following composition:—

	White Variety.	Red Variety.	Red Variety.
Water	75.50	75.20	69.10
Albuminoids	1.02	0.92	0.84
Fat	0.29	0.26	0.39
Sugar	5.19	5.82	8.42
Starch, &c.	15.52	15.13	15.81
Fibre	1.39	1.32	4.37
Ash	1.09	1.35	1.07

Large Radish.—One of the principal root crops in Japan consists of a giant radish, *Raphanus sativus*. No. 1 root was 30 inches in length, and weighed 3.65 kilos.; No. 2 was rather smaller:—

	No. 1.	No. 2.
Water	94.97	94.45
Albuminoids	0.57	0.64
Soluble carbo-hydrates	3.25	4.33
Fibre	0.60	0.58
Ash	0.61	0.58

Bamboo Shoots.—The underground buds of the root stock of the bamboo are frequently employed as food; they also serve for making pickles. Two principal kinds had the following composition:—

	Large variety.	Small variety (more mature).
Water	90.21	91.79
Albuminoids	3.28	2.59
Fat	0.13	0.11
Sugar	1.93	0.10
Extra active matter	2.54	2.81
Fibre	0.90	1.10
Ash	1.01	1.10

Seaweeds.—A list is furnished of a large number of edible seaweeds, and analyses of many of them are given. Laver (*Porphyra vulgaris*) is cultivated in the neighbourhood of Tokio. Branches of oak are placed in the shallow water of the bay in spring-time; on these the laver appears, and is collected from October to the following March, after which the plant becomes too hard to be of use. This seaweed flourishes best in brackish water. With the following analyses of Laver the market price of each is mentioned; it will be seen to follow pretty closely the percentage of nitrogenous matter, which is greatest in the young plant.

	Best.	Medium.	Common.	Purple.	Pale Green.
	Price 36 sen.	Price 29 sen.	Price 3 sen.	Price 27 sen.	Price 18 sen.
Water	14.40	12.60	19.40	12.98	12.91
Nitrogenous substance	26.14	18.11	4.48	17.41	19.88
Ext'd. matter	44.51	56.83	57.71	51.10	48.59
Fibre	5.50	5.66	7.46	9.83	9.98
Ash	9.45	6.80	11.90	8.68	8.64

100 Parts of the Ash contained—

Potash	34.50	31.50	11.15	35.19	33.83
Phosphoric acid	14.07	13.77	6.05	13.27	14.16
Silica	1.40	0.60	7.80	6.40	6.65

Analyses of Tangle (*Fucus saccharina*), and of other edible seaweeds, are given below. From the *Galidim corneum* the Kanten, or vegetable isinglass, is prepared. The cleaned plant is boiled with water, the solution is strained, and allowed to set to a jelly in wooden boxes. The jelly is cut into long prisms, frozen, and then allowed to thaw in the sun. The water runs away as thawing proceeds, leaving a white skeleton of kanten. One part will make a firm jelly with 150 of water.

* The albuminoids in all these analyses are found by multiplying the nitrogen by 6.33.

	Enteromorpha compressa.	Capea elongata.	Cystoseira sp.	Alaria pinnatifolia.	Fucus saccharina.	Vegetable Isinglass.
Water	13'60	13'17	16'40	15'11	26'80	22'80
Nitrogenous substance ..	12'41	8'99	8'42	8'29	7'79	11'71
Extractive matter	52'99	45'09	41'92	40'62	33'58	45'66
Fibre	10'58	7'40	17'06	2'16	9'33	4'97
Ash	10'42	24'74	16'20	33'82	22'50	18'53

100 Parts of the Ash contained.

Potash	27'98	—	32'55	21'00	27'00	31'77	—
Phosphoric acid — ..	11'22	2'37	2'20	2'61	4'43	2'96	—
Silica	6'97	2'20	1'91	trace	3'94	trace	—

Tea.—Three analyses of Japanese tea are given, two prepared by native methods, and one after the Chinese plan. 100 parts of tea contained:—

	Hiki-cha.	Sen-cha.	Chinese preparation.
Water	6'74	6'10	8'92
Soluble in Water ..	43'26	52'55	36'50
Tannin	12'50	12'10	13'19
Fibre	11'20	11'70	—
Ash	6'53	6'10	5'26
Nitrogen	5'79	6'38	3'18

Chrysalises.—The chrysalises of silkworms are used as food for fish, and also as manure. Two kinds contained as under:—

	Bombyx Mori.	Bombyx Yama-mai.
Water	10'99	9'28
Nitrogenous substance ..	47'28	49'75
Fat	14'83	23'57
Ash	3'24	2'54

Starch and Sugar.—Starch is prepared in Japan from buck-wheat; from the common brake, *P. aquilina*; from the root of a leguminous plant, *Pueraria Thunbergiana*; and from the root of a lily, *Erythronium grandiflorum*.

Starch-sugar is prepared by treating steamed rice with a portion of rice, wheat, or barley which has already undergone fermentation. After digestion for several hours at a fixed temperature the solution is strained, and evaporated either to a syrup or to a solid condition. The solid sugar contains 7 to 11 per cent of water. Both maltose and dextrin are present.

Alcoholic Liquors.—Rice-beer (*Sake*) is the most important beverage of the country. It is manufactured only during the winter months. The ferment is first prepared by spreading steamed rice mixed with wood-ashes in shallow trays, which are then seeded with the yellow spores of the fungus developed in a previous operation, and kept for some days in a warm chamber. When the fungus has developed, the mass is used to inoculate a fresh quantity of steamed rice. The mash is prepared by adding to steamed rice and water a certain proportion of the above ferment, and the whole is repeatedly agitated for two days. In this stage of the operation the starch of the rice is converted into sugar. The mash is then transferred to a large vessel, and the temperature raised to that of the human body. In this stage an active alcoholic fermentation takes place. The liquor thus formed is not, however, the final product. To it are added fresh quantities of steamed rice, water, and ferment, and after three days mashing this addition is again, and yet again repeated. When the last rice has been added the whole is allowed to ferment for thirteen days. The *sake* is now strained, and the residue pressed. After standing fifteen days to clear, the pure *sake* is heated to 120° F. to destroy any remaining yeast, and is then fit for use.

Finished *sake* contains, according to six analyses given by Mr. Kinch, 11'33 to 15'0 per cent of alcohol by weight, and 2'33 to 3'05 per cent of solid matter; the total free acid, reckoned as acetic, was 0'20 to 0'27 per cent. Various sweet, imperfectly fermented, liquors are also manufac-

tured. The solid residue left after pressing out the *sake* is distilled, and yields spirit of various strengths. The strongest spirit exhibited was 10'7 over Proof.

After the description of Japanese foods comes a summary of the dyestuffs used. Many of these are rapidly giving way to the brighter aniline colours.

The methods of preparing indigo and safflower are given in some detail. The six specimens of indigo exhibited contained from 10'91 to 19'3 per cent of pure indigotin.

The various oils and waxes of Japan form the concluding section of the catalogue. Vegetable wax is largely used for candles; it is prepared from the berries of the *Rhus succedanea*. The berries are collected in autumn, dried a few days in the sun, and then stored for several years. When taken for use they are crushed, steamed, and pressed while hot: a dark green wax is thus obtained, amounting to 15 per cent of the original berry. The wax may afterwards be purified and bleached.

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The determination of magnesium and its separation from the alkalies is effected by adding to the aqueous solution, whose volume should be about 25 c.c., a hot saturated solution of ammonium oxalate, or by dissolving in it solid ammonium oxalate with the aid of heat; boiling, adding an equal volume of acetic acid at 80 per cent, maintaining the mixture at a boil for some minutes with constant stirring, allowing to stand for six hours at 50°, filtering, and washing the precipitate with a mixture of equal volumes acetic acid, alcohol, and water. The precipitate and filter, still moist, are heated to dryness in a covered crucible, at first very gently, then with access of air, but with a small flame till the carbon is consumed, and finally strongly ignited, when the magnesium oxide is weighed. The latter is dense and heavy, and no loss is to be apprehended on ignition. In order to separate magnesium from the alkalies the oxalate must be precipitated from more dilute solutions, otherwise a double oxalate of magnesium and potassium is formed and is not decomposed by the subsequent treatment, whence the magnesium oxide is contaminated with potassium carbonate. The liquid, about 50 c.c., is therefore mixed with a cold saturated solution of ammonium oxalate (1:24), and $\frac{1}{2}$ vol. of alcohol is added. The magnesia thus prepared contains no potassium, or mere spectroscopic traces, and the precipitation of the magnesium is complete. The process is applicable whether the alkalies are present as chlorides, sulphates, or nitrates.

For the complete precipitation of manganese oxalate, Classen adds zinc chloride; in cases where the presence of

zinc would be inconvenient, he uses magnesium chloride, in such quantity that for each mol. of manganic oxide at least 1 mol. magnesia must be present, otherwise the residue after ignition is a mixture of manganic and manganoso-manganic oxides. For determining the zinc oxide the solutions are freed from all uncombined acid by evaporation in the water-bath, and in case of sulphuric acid by subsequent heating in the sand-bath; the residue is moistened with a few drops of dilute nitric acid, or digested for a short time with about 10 c.c. bromine water. A sufficient quantity of neutral potassium oxalate (1 : 3), about seven times the weight of the oxide, is added, the mixture is heated in the water-bath, and the ferric oxide is dissolved by the addition of acetic acid in drops. The whole is then heated to a boil, an equal volume at least of acetic acid is added, and after standing for six hours it is filtered while still warm, washed with the mixture of acetic acid and alcohol, and the zinc oxalate is ignited. Nickel and cobalt are determined in a similar manner; the washed oxalates are ignited, washed again, and re-ignited. The oxalate of nickel is precipitated in a crystalline state only when in small quantity, wherefore it is prudent to take only small quantities of the substance for analysis. Cobalt oxide is reduced in a current of hydrogen and weighed as metal.

Copper oxalate, if precipitated in the manner described for zinc and manganese, falls in a state of very fine division and settles badly. The determination is accurate and convenient if a sufficient quantity of potassium oxalate is added to the neutral concentrated solution. After a time the greater part of the copper separates in fine blue acicular crystals as cupro-potassic oxalate, and on adding an equal volume of acetic acid and allowing the liquid to stand, the residue of the copper is completely precipitated. The precipitate, after washing, is gently ignited, lixiviated, and re-ignited till the weight becomes constant, or, as Classen prefers, dissolved in dilute sulphuric acid, and the copper determined electrolytically by means of a thermobattery, or two Bunsen elements. In presence of antimony and arsenic chloride along with iron, as in the analysis of fahl-ores, the copper is not completely separated by the process above described, not even on the addition of zinc or magnesium chlorides. If but little antimony is present the substance is dissolved in nitric acid, the solution evaporated to dryness, the residue mixed with potassium oxalate in excess, filtered while hot, washed with a little water containing potassium oxalate, and the filtrate concentrated to 50 c.c. In presence of larger proportions of antimony the substance, finely pulverised and mixed with 4 parts ammonium chloride, is very gently ignited in a covered crucible, and thus almost all the arsenic and antimony, together with the greater part of the iron, are driven off as chlorides. The copper is then determined as above.

The separation of phosphoric acid from such oxides as form with potassium oxalate soluble double salts, capable of being decomposed by acetic acid, e.g., lime, is effected by the separation of the oxalates as described above. Ferric oxide and alumina, whose oxalates form with potassium oxalate double salts not decomposable by acetic acid, are completely eliminated by means of alcohol. The phosphate is dissolved in hydrochloric acid, evaporated to dryness, mixed with potassium oxalate six times the weight of the oxides, and digested for a short time in the water-bath, the ferric oxide dissolved by acetic acid and the same acid in excess and alcohol at 95 per cent are added as long as a precipitate is formed. After standing for six hours the precipitate is filtered, washed with alcohol, the filtrate in the beaker concentrated on the sand-bath to expel acetic acid and alcohol, the residue almost dry is diluted with water, any silica which separates is filtered off, and the phosphoric acid precipitated by means of ammonia and magnesia mixture. The process is suitable for the determination of phosphorus in crude iron. Arsenic acid behaves like phosphoric acid. Salts of cobalt, nickel, and zinc do not retain arsenic.

NOTES OF WORK BY STUDENTS OF PRACTICAL CHEMISTRY

IN THE LABORATORY OF THE UNIVERSITY OF VIRGINIA.

No. VIII.

Communicated by J. W. MALLETT,
Professor of General and Applied Chemistry in the University.

(Concluded from page 188.)

6. (51.) *Examination of the Constitution of Antimonic Acid prepared by Different Methods.* By C. P. CONRAD, of Winchester, Virginia.

According to Berzelius,* antimonious acid, obtained by acting upon metallic antimony with nitric acid or aqua regia, or by precipitating an antimoniate by an acid, has the composition $\text{Sb}_2\text{O}_3 \cdot \text{H}_2\text{O}$ (found 5.09 per cent H_2O , calculated 5.32 : atomic weight of Sb taken = 120).

According to Fremy† the product thus obtained is $\text{Sb}_2\text{O}_5 \cdot 5\text{H}_2\text{O}$ if dried in a stream of dry air at ordinary temperature (found 21.7 per cent H_2O , calculated 21.95), this being his antimonious acid; while he found his metantimonious acid, obtained by the action of water on antimony pentachloride, to consist of $\text{Sb}_2\text{O}_5 \cdot 4\text{H}_2\text{O}$ (found 17.1 per cent H_2O , calculated 18.37). More recently Geuther‡ states that by precipitating a solution of potassium antimoniate with nitric acid, washing the precipitate, and leaving it to itself during a whole summer, he obtained a hydrate having the composition $\text{Sb}_2\text{O}_5 \cdot 3\text{H}_2\text{O}$, which when heated to 175° gave off 2 molecules of water and became $\text{Sb}_2\text{O}_5 \cdot \text{H}_2\text{O}$. And Daubrowa§ says that by decomposing antimony pentachloride with water he prepared an antimonious acid which, dried in the air, had the composition $\text{Sb}_2\text{O}_5 \cdot 4\text{H}_2\text{O}$; dried over sulphuric acid, became $\text{Sb}_2\text{O}_5 \cdot 3\text{H}_2\text{O}$; at 100° became $\text{Sb}_2\text{O}_5 \cdot 2\text{H}_2\text{O}$; at 200° $\text{Sb}_2\text{O}_5 \cdot \text{H}_2\text{O}$; at 275° the anhydrous pentoxide, Sb_2O_5 ; and above 300° began to lose oxygen and become Sb_2O_4 .

In view of this discrepancy of statement as to the constitution of the antimonious hydrate or hydrates, Mr. Conrad undertook the experiments now reported.

Commercial metallic antimony, found to contain a small quantity of iron and traces of lead, tin, and copper, was purified by fusion with one-twelfth its weight of sulphur, oxidation of one-fifth the remaining metal by means of nitric acid, and fusion of the oxide so obtained with the other four-fifths of metal, this purification being repeated as often as was found necessary. The metallic buttons were cleaned on the outside by washing in acid and water.

From this pure metal antimonious acid was prepared by the following methods:—

- A.—Action of fuming nitric acid in excess, adding further amounts of acid several times, and evaporating finally on a water-bath until no trace of acid vapour could be detected by litmus paper.
- B.—Action of strong aqua regia, followed by evaporation with successive additions of strong nitric acid.
- C.—Decomposition of neutral potassium antimoniate (prepared as directed by Fremy) by hydrochloric acid.
- D.—Decomposition of acid potassium metantimonate (Fremy) by hydrochloric acid.
- E.—Decomposition of neutral potassium metantimonate (Fremy) by hydrochloric acid.
- F.—Decomposition of antimony pentachloride by excess of water.
- G.—Decomposition of antimony pentachloride by excess of a weak aqueous solution of ammonia.

* "Traité de Chimie" (Paris, 1830), ii., 499.

† Ann. de Chim. et de Phys., [3], xxiii., 404.

‡ "Jenaische Zeitschrift," vii., 121, quoted in "Watts's Dictionary," and Suppl., 98.

§ Ann. d. Chem. u. Pharm., 186, 110.

It appeared to be impossible to obtain by the first method (A) a product free from Sb_2O_4 . At any rate, in all the specimens actually prepared auric chloride revealed the presence of the lower oxide, and in one case a volumetric determination—by means of potassium pyrochromate, &c.—showed its amount to be equivalent to 20.86 per cent of Sb_2O_3 , or 44.04 per cent of Sb_2O_4 . No use, therefore, was made of the material prepared by (A).

Some of the products of (B), (F), and (G) also proved to contain antimonoso-antimonic oxide [Sb_2O_4 , or $\text{Sb}(\text{SbO}_4)$], to the formation of which there is clearly a very strong tendency; but specimens of antimonic acid quite free from this were made by all the processes save (A). In each case the product was very carefully washed, and the absence from the material taken for analysis of nitric acid, chlorine, potassium, and ammonium was established. In purifying (G) it was found necessary to wash at first with weak nitric acid.

In analysing the various specimens of antimonic acid, the antimony was determined (to avoid any uncertainty as to the character of the oxide present) in one portion by dissolving in a minimum of strong hydrochloric acid, adding tartaric acid, diluting largely, precipitating with hydro-sulphuric acid, heating for two or three hours while carbon dioxide was passed through the liquid, filtering on a weighed filter, drying to constant weight in a steam-bath, transferring most of the contents of the filter to a porcelain boat, placed in a hard glass tube through which a stream of carbon dioxide was passed, and gradually raising the temperature to redness so as to expel all traces of moisture and free sulphur; the true weight of the whole precipitate being calculated from that of the part thus purified. In a second portion of the same antimonic acid water was determined by exposure over sulphuric acid (in some instances at once in a steam-bath), and noting loss of weight; and afterwards by exposure to a slow stream of pure perfectly dry nitrogen at various determinate, progressively increased temperatures, the water driven off being collected in a chloride of calcium tube, and the loss of weight of the antimonic acid noted in comparison, so as to ascertain when reduction by heat to the lower oxide began. The remaining oxygen of the antimonic acid was obtained from the difference between the total weight and that of the antimony plus water. Preliminary experiments were made by this method upon pure, more or less hydrous, antimonious oxide with a view to practice, and to prove that trustworthy results could be obtained.

The loss of water was found to occur very gradually, and appeared to be influenced by the rapidity as well as temperature of heating. The results, however, point distinctly to $\text{Sb}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ as the composition of the acid dried in air at ordinary temperature and pressure over sulphuric acid, 14.53, 15.28, 14.45, and 14.11 per cent of water being found for 14.44 calculated. A larger, more variable, and more loosely retained (perhaps merely hygroscopic) amount of water was, in addition to this, found in the acid dried by simple exposure to air at mean temperature before placing it over sulphuric acid, making the largest amount found approximate to the $5\text{H}_2\text{O}$ of Fremy (21.16 to 19.12 per cent instead of 21.95 calculated).

For the samples dried at 100°C . the amount of water retained corresponded generally with $\text{Sb}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$: thus, 11.30, 11.44, 11.37, and 11.06 per cent were obtained instead of 10.11 calculated. This was verified by converting a known weight of pure metallic antimony into antimonic acid (by means of aqua regia), drying at 100° , and weighing. The result corresponded to 66.87 per cent of Sb in the product instead of 67.42 calculated for $\text{Sb}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$. But this point of temperature does not sharply mark the retention of this exact proportion, since figures as low as 8.95 and 8.77 per cent H_2O were in some cases observed. The temperature of commencing decomposition of $\text{Sb}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$ may fall below 100° .

Between 100° and 200° another molecule of water was given off, and $\text{Sb}_2\text{O}_5 \cdot \text{H}_2\text{O}$ formed; but it is clearly not easy to fix a precise temperature, at any rate without refer-

ence to other conditions, at which this hydrate either shall be obtained as a stable compound.

Finally, the statement of Daubrawa that the anhydrous pentoxide is produced at 275° was not verified. Even after prolonged exposure to a temperature of 300° as much as 2.74 per cent of water (about half a molecule) was found to be retained, and could only be completely expelled at a low red heat, when oxygen was also lost and Sb_2O_4 began to be formed.

The series of specimens examined was not quite complete, owing to one or two analyses having been lost, and not sufficient material being on hand for their repetition; but there was no indication of any inherent difference of composition in the acids prepared by the various methods, and Fremy's statement under this head in regard to free antimonic acid failed of verification.

On the whole, there seems to be sufficient reason for believing in the existence of three antimonic acids corresponding with the three chief hydroxylic acids of phosphorus, and it will conduce to clearness if the nomenclature be made to agree with that of these latter. Thus we shall have—

H_3SbO_4 , or $3\text{H}_2\text{O} \cdot \text{Sb}_2\text{O}_5$, Ortho-antimonic acid.

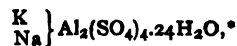
$\text{H}_4\text{Sb}_2\text{O}_7$, or $2\text{H}_2\text{O} \cdot \text{Sb}_2\text{O}_5$, Pyro-antimonic acid (Metantimonic acid of Fremy).

H SbO_3 , or $\text{H}_2\text{O} \cdot \text{Sb}_2\text{O}_5$, Metantimonic acid (antimonic acid of Fremy).

The so-called acid antimonates and metantimonates of Fremy, if not containing basic hydrogen, are of course not true acid, but anhydro-salts, compounds of the normal salts with antimony pentoxide. The gradual, rather than sudden, transition on heating from the first of these metallic acids to the second and third finds a parallel in the corresponding case of the acids of phosphorus; and the similarity of behaviour of the latter even extends in some degree to the final production of the anhydrous oxide, since H. Rose has observed that when meta-phosphoric acid is volatilised the last portions are mixed with phosphorus pentoxide. In the case of antimony there is a notable tendency, both in the production of the acids and in their decomposition by heat, to partial loss of oxygen and formation of ortho-antimonate of antimony, $\text{Sb}(\text{SbO}_4)$ or Sb_2O_4 , the well-known neutral oxide of the metal.

7. (52.) *On the Mutual Relations, in Aqueous Solution, of Potassium and Sodium Alums.* By F. P. VENABLE, of Charlottesville, Va.

Two forms of isomorphism between these salts are conceivable, viz., (a) the replacement in potassium alum of one out of the two atoms of potassium by sodium, or reciprocally as to sodium alum, giving rise to a salt of the constitution—



which might be confidently expected to exhibit similarity of crystallisation to the two alums with a single alkaline metal each; and (b) the isomorphous admixture, in various proportions, of the ordinary alums themselves in individual crystals. If the formation of the double alkaline alum were possible, such a salt would most probably be capable of crystallising in all proportions along with either simple potassium or sodium alum, but as these two latter differ from each other greatly in solubility it was thought possible that some indication might be obtained of a tendency to the formation of the potassio-sodium alum by crystallisation, effected under various conditions of temperature, strength of solution, and relative proportion of the two simpler salts.

With a view to test this, perfectly pure potassium and sodium alums were prepared, free from ammonium salt, and their solutions were mixed in proportions varying from

* It being assumed, on the strength of the experiments of Hertwig (Pogg. Ann., lv., 59), Lupton (Jour. Chem. Soc., [2], xiii., 201), and Heintz (Pogg. Ann., lv., 331) that $\text{M}_2\text{Al}_2(\text{SO}_4)_4 \cdot 24\text{H}_2\text{O}$, and not this formula halved, correctly represents the ordinary alums.

x molecule of sodium alum for 1 of potassium alum to 25 mols. of the former for 1 of the latter. The mixed liquids were exposed to spontaneous evaporation for different periods, and at temperature ranging from 5° to 25° C., but no tendency could be traced to the uniform production, under any given set of conditions, of crystals containing x atom potassium for 1 of sodium. The isomorphous replacement seemed to be solely of the second or simpler kind (b), while the inferior solubility of potassium alum tended strongly to make this salt relatively predominant in the crystals separated.

In the course of these experiments it became apparent that, as in some of the cases, examined by Karsten* and Page and Keightley,† of mixed solutions of salts not acting chemically upon each other, the amount of each dissolved was diminished by the presence of the other, and as no results of this kind in regard to the alums seem to be on record, it may be well to quote the following table, giving the figures obtained for the effect on the solubility of potassium alum produced by the presence of various amounts of sodium alum. The several solutions of the latter were placed in contact with an excess of finely-crystalline potassium alum in closed flasks, allowed to stand for four days, with occasional agitation, and the clear liquid then drawn off and analysed. The temperature ranged from 13° to 16° C. Supersaturation, it is believed, was effectually guarded against.

Composition of 100 c.c. of each Solution.

No. I.	Water. Grms.	Sodium Alum. Grms.	Potassium Alum. Grms.
" 2.	97.4	4.70	7.60
" 3.	94.9	9.46	5.75
" 4.	93.8	11.35	5.33
" 5.	92.3	14.19	4.85
" 6.	89.7	18.92	4.22
" 7.	84.1	28.38	3.23
" 8.	76.5	42.57	2.07
" 9.	74.0	56.76 (sat.)	1.28

Or, expressing these results otherwise, 100 grms. of water, if already holding in solution the amounts of sodium alum noted in col. 2 of the following table, will dissolve at most the amounts of potassium alum given in col. 3.

		100 grms. of Water, Sodium Alum { will { dissolve of } Potassium Alum	
No.		4·8 grms.	7·8 grms.
" 2.	" "	10·0 "	6·1 "
" 3.	" "	12·1 "	5·7 "
" 4.	" "	15·4 "	5·3 "
" 5.	" "	21·1 "	4·7 "
" 6.	" "	33·7 "	3·8 "
" 7.	" "	55·6 "	2·7 "
" 8.	" "	76·7 "	1·7 "

University of Virginia,
July 25, 1879.

ON A

POSSIBLE CAUSE OF VARIATION IN THE PROPORTION OF OXYGEN IN THE AIR.

By E. W. MORLEY, M.D., Ph.D.,
Professor of Chemistry in Western Reserve College, Hudson, Ohio.
(Concluded from page 186.)

WHEN the writer planned to make analyses of air in order to detect if possible some law in its variations of composition, he expected to have to do with quantities but little larger than the errors of observations. Some thought was therefore given to the methods of keeping such errors as small as possible. It was hoped that if the probable

error of a determination of oxygen with this apparatus was not larger than the probable error of the analyses made by Bunsen in January and February, 1846, the object of the analyses would be attained. Such analyses as those of Bunsen would abundantly serve to establish variations of the four hundredth part of the average amount of oxygen contained in the atmosphere. The writer expected to have commonly to do with such variations, and therefore computed the comparative accuracy of analyses of air made with the long eudiometer of Bunsen's experiments, and with the apparatus used in the present work. In the first analysis by Bunsen of the air of January 9th, the length of the column of gas in the eudiometer was in round numbers 840 millimetres, and its tension 570. The tension was determined by four readings, and the apparent volume of air taken by one reading. An error in the last reading would produce also an error of the same sign in the observed tension, so that these two errors are not independent. Their influence on the result is therefore computed by simply adding them; the other errors are independent of this joint error and of each other. And the influence of the four independent errors on the result is computed by taking the square root of the sum of their squares. If we repeat this computation for the different volumes and tensions of the second and third measurements, referring all the probable errors to the volume of air first taken, and obtain the probable errors of the three measurements, we may obtain the probable error of the final result by adding the square of the first probable error to the squares of the third parts of the second and third; and taking the square root of the sum.

The probable error of a single reading of the level of the mercury in Bunsen's experiments is not given; but we may compare the two methods by assuming arbitrarily some probable error, provided that we assume the same error in both computations. The first column of the following table gives the influence of a probable error of one-tenth of a millimetre in a single reading in each of the three measurements of the analysis quoted from Bunsen.

In the measurement of a volume of gas with the Frankland and Ward or McLeod apparatus, the mercury is brought to that mark in the eudiometer tube which will give a suitable tension, and the height in the pressure tube at which the mercury stands is determined. An error in determining the volume of gas in the eudiometer tube involves an error of the same sign in the observed tension. If we add these two dependent errors we have one of the two independent errors affecting this mode of measurement, the other being the error probably made in determining the upper level of the mercury in the pressure tube. Adding the squares of these two and taking the square root, we have the probable error of a measurement with the apparatus. If we treat the second and third measurement in the same way, and compute the effect of these three probable errors on the final result, we get the numbers in the second column of the following table.

*Probable errors of measurements of gas, and of final results,
occasioned by a probable error of the tenth of a
millimetre in each reading.*

	In Analysis cited from Bunsen, Per cent.	With Frank- land and Ward apparatus. Per cent.
In measurement of air taken	0'046	0'034
In measurement of air and hydrogen ..	0'050	0'042
In measurement after explosion	0'039	0'031
Probable error of result	0'051	0'038

It is obvious that with the same error probable in each reading, the use of the rapidly working apparatus involves no sacrifice of accuracy to convenience, as far as the conditions of *observation* are concerned.

To obtain the degree of accuracy needed for the present purpose, it is necessary to take account of the expansion of the mercury in the column which measures the pressure,

* *Schrift d. Berl. Akad.*, 1841.

† *Jour. Chem. Soc.*, [2], x., 566.

and also of the linear expansion of the scale which measures the tension, and of the cubical expansion of the eudiometer tube. Since all these are at the same temperature with the gas to be measured, it is easy to prepare a table giving the correction not only for the expansion of the gas but also for the expansion of mercury, scale and eudiometer. Such a table, giving on a single page, for every tenth of a centigrade degree from zero to thirty degrees, the logarithmic factor to be added to the logarithms of the observed volume and observed tension, makes the work of reduction very slight. Most tables of correction of the volumes of gas contain a logarithm to be subtracted; for convenience it should be additive; and five places of decimals should not be exceeded. Unless measurements can be made ten times more accurate than Bunsen's, five places of logarithms distinguish smaller differences than observation deals with; five places permit instant interpolation for hundredths of a degree, and a greater number waste time and possess no advantage whatever.

The eudiometer was calibrated by filling it with air-free water, and weighing the quantity expelled as the mercury rose to each successive mark of the graduation. This was done four times for each division; the probable error of a single determination was found to be 8.6 milligrammes of water. The results were all reduced to the temperature of melting ice. There were seventeen divisions in the eudiometer tube, now broken, which was used in all the analyses in this paper. The volume of gas to be measured was always brought to one of the two divisions which permitted measurement under the most favourable conditions, and its tension determined; it was then brought to the other division, and its tension again determined. Two independent measurements thus obtained eliminated the chance of error in identifying divisions on the scale, and also afforded the means of ascertaining the probable error of a measurement. In the analyses contained in this paper, the quantity of air taken was unfortunately limited by the circumstance that the collecting tubes at hand were small; the probable error of the results so far obtained is therefore much greater than is due to the care used in observing. In analyses made after the present month this mistake of judgment will be corrected. The mean quantity of air taken in an analysis so far has been 38.9 c.c. measured at zero and 760 millimetres. From 196 pairs of measurements it has been computed that the probable error of a single determination of volume is its 5800th part. Hence the probable error of a determination of oxygen in the air is the 7200th part, and the probable difference of two determinations on the same sample is the 5100th part. A second analysis was always made when the first showed a deficiency of oxygen. A comparison of the results will show whether the accuracy indicated by computation was obtained.* The writer has in hand an entirely new construction of the pressure tube, and some modifications of the optical appliances for reading the level of the mercury in the eudiometer tube, by which he hopes considerably to lessen this probable error.

The samples of air analysed were collected in the open country in glass vessels with due care as to admixture with the air from the collector's lungs, preserved over mercury, freed from carbonic acid, and exploded with hydrogen, of proved purity, obtained by galvanic decomposition of water. But some samples were collected in clean stoppered and capped bottles, and kept for a short time by inverting the bottle in the cap which had been filled with water. In this case the air was withdrawn for analysis with a Toepler's mercury pump.

The table gives in the first column the date, in the second the mean temperature of the day at this place as determined by three observations. In the third, on the

days when analyses were made, the hour of collecting the sample is given, fractions of hours being disregarded. In the last two columns are given the hundredths per cent found by analysis, the figures twenty and the decimal point being suppressed. The figures ninety-six, for instance, in this column mean 20.96 per cent of oxygen. Within the time covered by the analyses now published, there were several well marked great and sudden depressions of temperature, and the figures show the falling off in the proportion of oxygen in the air at these times to be as well marked as the depression of temperature. The deficiency is not proportionate to the depression of temperature; this could not be expected.

ANALYSES OF AIR,

Showing Deficiency of Oxygen attending Sudden Depression of Temperature.

Winter of 1878-1879.

Date.	Mean Temperature. F.	Hour of taking sample.	Oxygen.
Dec. 28	—	4 P.M.	98 96
29	14.8		
30	19.3		
31	16.2		
Jan. 1	23.8		
2	7.6	4 P.M.	91 92
"		10 P.M.	90 89
3	-7.2	9 A.M.	90 91
"		1 P.M.	96 97
4	2.2		
5	7.7		
6	13.5	3 P.M.	97
7	19.3		
8	26.4		
9	16.9		
10	9.6	10 A.M.	96
11	20.6		
12	26.0		
13	27.9		
14	26.3		
15	25.8		
16	28.5		
17	29.1		
18	—		
19	15.9		
20	9.9		
21	26.0		
22	37.7		
23	28.3		
24	32.5		
25	32.4		
26	23.7		
27	44.9		
28	37.4	9 A.M.	96
29	31.5		
30	—		
31	28.2		
Feb. 1	18.5	9 A.M.	96
"		9 P.M.	94 94
2	19.8	9 A.M.	91 93
"		9 P.M.	82 80
3	26.8		
4	28.1		
5	31.5		
6	28.3		
7	25.1		
8	28.5		
9	24.0		
10	32.7		
11	54.4		
12	26.5		
13	15.6		
14	5.8		
15	11.1		

* The divergence of the second result of February 26th from the first is due to the fact that in the second analysis the hydrogen used was not pure. As none of the sample remained for a third experiment, the second result is given in confirmation of the first. But this pair of results should not be used in computing probable errors.

Date.	Mean Temperature. F.	Hour of taking sample.	Oxygen.
Feb. 16	26.3	9 A.M.	95
17	25.7		
18	24.1		
19	23.2		
20	18.9	6 P.M.	87 87
21	17.9		
22	—		
23	25.4		
24	21.5		
25	37.1		
26	22.3	3 P.M.	45 50
27	12.8	9 P.M.	77 80
28	24.1		
Mar. 1	38.7		
2	29.7		
3	34.3		
4	37.2		
5	35.9		
6	41.6		
7	34.1		
8	51.0		
9	61.8		
10	58.4		
11	54.4		
12	40.3		
13	39.4		
14	29.3		
15	22.8	9 A.M.	88 84
"		9 P.M.	84 86
16	25.3	9 P.M.	92 92
17	24.5	9 A.M.	89 90
18	24.3		
19	28.7		
20	33.1		
21	32.3		
22	34.6		
23	31.7		
24	40.1		
25	30.1		
26	38.2		
27	35.1		
28	44.8		
29	45.9		
30	33.3		
31	31.9		
Apr. 1	33.7		
2	29.9		
3	{ 25 at }	9 A.M.	77 79
"	{ 2 P.M. }	9 P.M.	85 87
4	27.1	9 A.M.	80 80
"		9 P.M.	88 85
5	28.2	9 A.M.	77 77
"		9 P.M.	86 82
6	39.4		

It may be said that these analyses were commenced in March, 1878, but in December of that year, a doubt was felt whether it was absolutely certain that in every case the air and hydrogen had been completely mixed before explosion. In test cases, the air and hydrogen had been permitted to diffuse into each other for eighteen hours before explosion, and the results were the same as in the usual course of analysis; but the analyses are not here given, although they contain only evidence perfectly agreeing with that here presented. In all the analyses here printed, the air and hydrogen were known to be thoroughly mixed; they were driven as rapidly as possible through a capillary tube twelve or twenty times. All made between the first and last dates of the table are given without selection, except that some were rejected for obvious instrumental errors.

The remarkable deficiency of oxygen observed on the 26th of February seems affected with no reason for doubt. On Sept. 16, 1878, two very careful analyses

of the same sample gave 20.49 and 20.46 per cent of oxygen. On July 19, and Nov. 10, 1877, Jolly found 20.56 per cent. The *Neues Handwörterbuch der Chemie*, i., 856, cites an analysis of air from the Bay of Bengal showing 20.46 per cent, one of air from near Calcutta, showing 20.39 per cent, and one of air from near Algiers, showing 20.41 per cent. That Jolly and the writer have found air almost as deficient in oxygen as the three last will lessen the probability that the air of the surface of the earth in the Torrid zone is normally poor in oxygen. One of the first cases of a supposed descent of cold air from an elevation mentioned by Loomis occurred in the warmer parts of this country. If his theory finds favour, and the writer's conjecture is correct, it will be presumed that the three samples cited in the *Handwörterbuch* from the still warmer regions of the earth were taken in the midst of such a mass of cold air descending from, and retaining the composition of, the upper part of the earth's atmosphere.

The analyses here printed should not be used in determining the average composition of the air by combining analyses from all sources. Whether the writer's conjecture is correct or not, it has enabled him to select times for taking samples of air varying widely from the average; and to such times his analyses have been commonly limited, only occasionally including a sample of presumably normal air to serve as a check on the abnormal.—*American Journal of Science*, September, 1879.

NOTICES OF BOOKS.

Friction and Lubrication. Determination of the Laws and Coefficients of Friction by New Methods and with New Apparatus. By R. H. THURSTON, Professor of Mechanical Engineering at the Stevens Institute of Technology. London: Trübner and Co.

THIS work, which bears as its second title the words "Lectures on Friction, Lubrication, and Lubricants," seems to be a very complete and useful monograph on an important subject. Our attention is naturally in the first place drawn to what may be called the chemical department of the treatise,—the account given of the various lubricants, of the properties on which their applicability depends, and of the methods by which their applicability is tested. This amounts substantially to an account of oils, their sophistications, and their commercial analysis. For an oil to be really trustworthy as a lubricant it must combine a number of properties; it should be not liable to ignition or decomposition at any temperature likely to be produced by friction; it must have no corrosive action upon metals, and should therefore be free from acidity; it should not clog or "gum" on prolonged exposure to the air—a requirement which excludes all the so-called "drying" oils; it must not be liable to congeal on exposure to cold; it should minimise friction and should possess a maximum power of carrying away and dissipating heat, so as to keep cool the surfaces to which it is applied. These are general properties required for all lubricants. The degrees of body and of fluidity preferred vary according to the character of the machinery in question. Some of these qualities are estimated by chemical experiments, whilst others can only be judged by mechanical treatment approaching as closely as possible to the conditions of actual practice. The genuine or spurious nature of a given sample of oil is another consideration on which Professor Thurston bestows no little attention. Here it must be remembered that the question of purity can only arise if an oil is sold under a name denoting its origin. If a dealer offers us, e.g., olive oil, we have a right to object if the sample contains even a trace of an oil from any other source, and that quite irrespective of any appreciable deterioration arising from such admixture. But if the

sample is offered merely as a lubricating oil, then, so long as its beneficial action is in fair proportion to its market-price, the purchaser suffers no wrong.

We fear that the methods here given for detecting the various constituents of a mixed oil, except in thoroughly experienced hands, will often fail of arriving at the whole truth. Such mixtures are often somewhat complex, and contain not merely two or three kinds of oil, but other substances in solution, some of them added for the express purpose of masking reactions. Again, not only do the chemical characters of any given oil vary with its age, with the country where the raw material has been grown, and with the manner of its manufacture, as the author fully recognises. It seems probable that the reactions by which we distinguished one oil from another depend not so much on the oils themselves as on the presence of certain naturally accompanying impurities. In proportion as these are got rid of by refining, the distinctions upon which the analyst must rely fade away.

The rise of temperature produced by different oils on the addition of concentrated sulphuric acid has been proposed as a distinguishing test both by Maumené and Fehling. But the results of these two chemists are utterly irreconcilable. Thus, according to Fehling the four oils linseed, sesame, castor, and cod-liver, all show an increase of temperature of 74° (Fahr. or Cent. ?), whilst Maumené gives the respective rise as 133°, 68°, 47°, and 103°. No one can use such a process until it has undergone a careful revision in competent hands. Chateau's process for the examination of fatty bodies as reported in the *Bulletin de la Société Industrielle de Mulhouse*, 1861, is quoted at some length, as is also the "oleography" of Prof. Tomlinson and Dr. Moffat (see *CHEMICAL NEWS*, vol. xviii., p. 299), which, to be satisfactory, requires great care and nicety of manipulation.

The method which the author recommends for determining the liability of any particular oil to gumming and drying is that of Nasmyth, and consists in observing how far a drop of the oil will travel in a groove cut in an inclined plane of iron or glass. We should strongly advise consumers of lubricating oils not to attempt their chemical examination in person, but commit it to a professional analyst.

The work before us is provided with a very copious index, but we regret to perceive a great number of typographical errors.

Structure and Development of the Brain. A Lecture delivered in the City Hall, Glasgow, under the auspices of the Glasgow Science Lectures Association. By ALLEN THOMSON, M.D., F.R.S. London and Glasgow: W. Collins, Sons, and Co.

WE have here an account of the brain, clear, compact, and accurate; embodying the results of the latest investigations, and rendered perfectly intelligible to the non-professional mind. The learned lecturer deals successively with the origin, early formation, and development of the brain in the embryo, and the differences in its form, size, and structure among animals. He compares the human brain with that of the anthropomorphous apes, and touches on the varieties of form and size observed in the brains of different races and individuals. Lastly, he traces the relation and connection of the various portions of the brain, having due regard not merely to its more obvious structure, but to its microscopical texture.

Dr. Thomson fully admits that the differences of form and structure observable in the brains of the vertebrate animals may be regarded in some measure as a repetition of those presented by the embryo brains of the higher animals in the successive phases of their development, phylogenesis and ontogenesis being thus parallel. But he points out that in the convoluted condition of the cerebral hemispheres there is no regularly increasing gradation. Nearly smooth and highly convoluted brains

may be met with, even within one and the same mammalian order. Thus, among the Monotremes, the brain of *Ornithorynchus* is nearly smooth, whilst that of *Echidna* is strongly convoluted. Even in the Primates the plane hemispheres of the *Hapalida*, such as the Marmoset, contrast strikingly with the highly developed convolutions in man and in the anthropoid apes.

The relation of these convolutions to intelligence will appear at least questionable if we reflect that in the Rodents they are very small in size. Yet the intelligence of at least one rodent form, the common rat, is but too well known.

In comparing the brains of the European races with those of savages, the author seems to overlook the perplexing fact that the average Esquimaux brain is at least equal in size and weight to that of civilised nations. The weight of the brain in relation to that of the whole body fails likewise to throw any satisfactory light upon the gradation of intelligence. The brains of the smaller monkeys and of certain birds, such as finches, are proportionately double the weight of that of man.

When we add, in conclusion, that this treatise can be procured for the sum of three pence, the ignorance concerning the structure and functions of the brain, which is met with even among people of culture, must be considered simply inexcusable.

General Physiology of the Nervous System. A Lecture delivered in the City Hall, Glasgow, under the auspices of the Glasgow Science Lectures Association. By J. GRAY M'KENDRICK, M.D., &c. London and Glasgow: W. Collins, Sons, and Co.

THIS lecture may be considered as a continuation of that of Dr. Allen Thomson. The author reviews the entire nervous system not so much from a structural as from a functional point of view. The author explains the points of similarity, and also the distinctions between electrical conduction and nervous action, and shows that in man, and probably in other warm-blooded animals, the propagation of the latter is more than a thousand times slower than the speed of electricity along a copper wire. The nature of reflex movements is next explained—a subject on which much popular misapprehension prevails, and is eagerly utilised by humanitarians and sentimentalists of the day. The functions of different portions of the brain are next discussed, as far as our very rudimentary knowledge will allow. Lastly, the author turns to the connection between mental phenomena and the varying physical states of the brain. Admitting the existence and the intimate character of such connection, he very judiciously declares that we have no evidence for regarding the mental states as simply the product of such physical changes, or for assuming that when these physical operations cease the mental states cease also.

These "Science Lectures," if the attendance has been in any degree proportionate to their merit, must have been productive of great good, which will be further heightened by their re-appearance in the present cheap and convenient form.

CORRESPONDENCE.

PRESENCE OF ALCOHOL IN ANIMAL TISSUES DURING LIFE.

To the Editor of the Chemical News.

SIR,—Years ago the late Dr. Anstie and myself showed that animal tissues, when distilled with water, yield a distillate containing traces of a substance which gives all the characteristic reactions of alcohol. In our researches

on the elimination of alcohol, we had not only recognised this, but made allowance for the presence of this natural alcohol. The discovery is, therefore, not a recent one, and is not due to T. Béchamp.—I am, &c.,

A. DUPRÉ.

October 18, 1879.

METHOD FOR THE ESTIMATION OF THE OXIDE OF IRON AND ALUMINA IN COMMERCIAL PHOSPHATES.

To the Editor of the Chemical News.

SIR,—The following method of estimating the oxide of iron and alumina existing in phosphates, which I have found very expeditious, and more reliable than the ordinary process of precipitating with acetate of ammonium, may be found useful by some of your readers.

Determine the total P_2O_5 in the sample by precipitation with magnesia mixture. Dissolve a fresh portion of the sample in HCl plus a few drops of HNO_3 , precipitate the lime as oxalate without troubling to separate the silica, filter, add a few crystals of potassium chlorate to the filtrate, boil well, add a slight excess of ammonia, boil, and filter off the precipitate of iron and alumina phosphates, dry, burn, and weigh it. In the filtrate estimate the P_2O_5 remaining in solution by standard uranium; the amount found subtracted from the total amount in the sample gives the P_2O_5 in the iron and alumina precipitate; subtracting this, therefore, from the weight of the precipitate, we get the oxide of iron and alumina present in the sample. The method does not take long and gives very accurate results.—I am, &c.,

T. TWYNAM.

Chemical Laboratory and Assay Offices,
106, Leadenhall St., London, E.C.,
October 20, 1879.

PYRITES-SMALLS IN THE MANUFACTURE OF SULPHURIC ACID.

To the Editor of the Chemical News.

SIR,—Since that matter is of considerable interest for British vitriol manufacturers, I beg permission to add a few words to the remarks made by Mr. Watson Smith in the CHEMICAL NEWS, vol. xl., p. 189, on the burning of pyrites-smalls. The amount of sulphur left in the burnt ore, where the "shelf-burner" is used, is not 5 per cent, but never above 1½, sometimes only ½, on the average perhaps 1 per cent. This I found to be the case both in France and in Germany, as well as at the Uetipon Works, near Zürich, referred to by Mr. Smith, whose description is perfectly correct in every other way. The raw material used here and in France is 48 per cent Chessy pyrites, in North Germany 44 per cent Schwelm pyrites. The burnt ore is taken straight from the vitriol works to blast-furnaces, being pure enough for this purpose. It is strange that this most efficient of all pyrites-burners is so little known or appreciated in England, although (or perhaps because) it is not protected by a patent. In my "Treatise on the Manufacture of Sulphuric Acid and Alkali," there are diagrams both of the original Malétra burner and of the improved Aussig burner, of which my friend M. Schaffner has enabled me to give the fullest working drawings. The Aussig burners are immediately connected with a Glover tower, which proves that the difficulty caused by flue dust has been entirely overcome in their construction; and both the yield of vitriol and the consumption of nitre in that place are among the most favourable I ever heard of.—I am, &c.,

GEORGE LUNGE.

The Polytechnicum, Zürich,
October 20, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 14, October 6, 1879.

Artificial Laurite and Ferriferous Platinum.—H. Sainte-Claire Deville and H. Debray.—The common metals, such as iron and lead, exist in nature in various combinations, but the most abundant which constitute their true ores are precisely those which we see formed under her eyes, when these same metals are abandoned to the action of atmospheric agencies. Laurite may be obtained by heating a mixture of ruthenium and iron pyrites. The sulphur resulting from the decomposition of the pyrites combines with the ruthenium, and this sulphide dissolves in the ferrous sulphide, and crystallises on cooling in regular octahedra, like natural laurite. All the sulphides of the platinum metals are decomposable by heat at a temperature sufficiently high. By heating laurite until the crucible began to soften, metallic ruthenium was obtained in small cubic crystals. A platinum sulphide was obtained by melting platinum with ten times its weight of pyrites, no compound similar to laurite being formed. If the above mixture is very strongly heated, the regulus, after treatment with hydrochloric and nitric acid and potassa, yields ferriferous platinum.

A Sporadosideric Meteorite, which fell January 30, 1879, at La Bécasse, Indre.—M. Daubrée.—The meteorite is 2'800 kilos. in weight; its paste or main mass consists of peridot and a bisilicate, like pyroxene or enstatite, interspersed with metallic grains of nickeliferous iron, accompanied by troilite.

Mathematical Theory of the Changes of Brightness of Variable Stars.—H. Gylden.—Not suitable for abstraction.

Synthesis of a Diphenyl-propane and on a New Method of Formation of Dibenzyl.—R. D. Silva.—On causing ordinary propylene chloride to act upon benzol in presence of aluminium chloride, the author obtained a liquid of an agreeable odour, and boiling without decomposition at the atmospheric pressure, which he supposed to be a diphenyl-propane. On endeavouring to produce allyl-benzol by causing chloride of allyl to act upon benzol in presence of aluminium chloride, he obtained diphenyl-propane.

Reaction of Cyanamide upon Hydrochlorate of Dimethyl-amin.—P. Tatarinoff.—Dimethyl-guanidin is formed by the action of heat upon cyanamide and dimethyl-amin hydrochlorate in molecular proportions. The composition of its platinum compound is— $(C_3H_9N_3, HCl)_2PtCl_4$.

Mineral Associations Contained by Certain Trachytes of the Riveau Grand, Mont Dore.—F. Gonnard.—Not suitable for abstraction.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin No. 9, 1879.

Atomic Weight of Antimony.—F. Kessler.—The author criticises the researches of J. P. Cooke and considers that the atomic weight of antimony may still be taken at 122.3 (O = 16), or 122 (H = 1, O = 15.96).

Liquid Toluol-sulpho-chloride, and the Toluol-meta-sulphuric Acid of Beckurts.—C. Fahlberg.—The author considers it indubitable that the toluol-meta-sulphamid of Beckurts consists of a mixture of toluol-para-sulphamid and toluol-ortho-sulphamid.

Benzal-sulphide and Sulpho-benzaldehyd.—C. Böttinger.—Not suitable for abstraction.

Behaviour of the Acids of Nitrogen with Sulphuric Acid.—G. Lunge.—Hyponitric acid cannot under circumstances exist as such in contact with sulphuric acid, but is decomposed into nitrous acid, which immediately forms nitrosyl-sulphuric acid (chamber crystals) with a portion of the sulphuric acid, and nitric acid which dissolves as such. The nitrosyl-sulphuric acid dissolves in the excess of sulphuric acid, forming a colourless liquid, but only up to a certain limit, which may be regarded as the saturation-capacity of sulphuric acid for nitrosyl-sulphuric acid, and which rises with the concentration of the sulphuric acid. Above this limit there is produced a yellow colouration. As such acids become colourless on prolonged boiling, the excess of nitrosyl-sulphuric acid appears to be in an unstable state of combination; it does not, however, appear affected by the temperature of the water-bath. The circumstance observed by Winkler that a mixture of sulphuric acid at 66° B. and hyponitric acid displayed an orange colour, emitted red vapours, and that a violent disengagement of hyponitric acid took place on the application of heat, proving that this latter acid was present unchanged, can only occur if the mixture contains far more hyponitric acid than ever occurs in the manufacture of sulphuric acid. Every nitrose (*i.e.*, solution of nitrosyl-sulphuric acid in sulphuric acid), whether nitric acid is also present or not, takes, if heated far below its boiling-point, a golden or still darker yellow shade, but becomes colourless again on cooling. This change of colour can be repeated at pleasure; it points scarcely to a loosening of the compound, which displays great permanence even at a much higher temperature, but may rather be compared with the heightening of the colour of ferric chloride when heated. The stability of nitrosyl-sulphuric acid dissolved in sulphuric acid is great, even at the boiling-point of the latter if its sp. gr. is not below 1.70. Nitrogen is indeed continually lost during boiling, and the more the less concentrated is the acid. But if the boiling is so conducted that the escaping vapours cannot be condensed and flow back, nitrosyl-sulphuric acid is found in the residue, even at sp. gr. 1.65°. If the vapours are condensed and flow back, a considerable loss takes place by denitration. Down to 1.65 sp. gr. the affinity for nitrous acid, *i.e.*, the tendency to form nitrosyl-sulphuric acid, is so great that nitric acid simultaneously present, whether added as such or formed by the decomposition of hyponitric acid, is reduced with escape of oxygen, and is expended in the formation of nitrous sulphuric acid. With acid of sp. gr. 1.71 or still stronger, this conversion is almost complete after a short boiling; at 1.65 sp. gr. it is incomplete. This is a further argument against the presence of hyponitric acid in the solution. Below 1.65 sp. gr. nitrosyl-sulphuric acid is so unstable that most of it, if not all, is expelled from an acid of 1.6 sp. gr. by a short boiling. In an acid of sp. gr. 1.5 partial decomposition of the nitrous acid occurs, even in the cold, with formation of nitric acid and nitric oxide, but after heating for an hour in the water-bath considerable quantities of nitrosyl-sulphuric acid remain undecomposed, whilst another part passes into nitric acid. On boiling the former portion is totally lost. In consequence of the very low saturation-capacity of sulphuric acid at 1.5 sp. gr. for nitrosyl-sulphuric acid, free nitrous acid is present along with the latter in the cold. In still weaker acids this occurs in a higher degree; it is, however, probable that even very dilute sulphuric acid can contain, in the cold, a little nitrosyl-sulphuric acid if reducing agencies are absent. The nitric acid co-existing with nitrosyl-sulphuric acid remains in great part in the liquid after long boiling, and even in acids of sp. gr. 1.5 and lower. If, therefore, the nitrose from the Gay-Lussac tower of a vitriol works in consequence of bad management contains nitric as well as nitrous acid, it cannot be completely denitrised by hot water or steam, not going below 48° B. = 1.5 sp. gr. The end can only be effected by reducing agents, as on the large scale by sulphurous acid in the Glover tower, and on the small scale by mercury in the

nitrometer. It can be plainly seen in the latter how much more difficult and tedious the process of denitrification becomes if nitric acid is present. The tendency to the formation of nitrosyl-sulphuric acid is so strong that even when there is a very large simultaneous introduction of air and consequently of oxygen into the sulphurous acid, along with nitrous acid or nitric oxide, no oxidation to N_2O_4 or NO_3H can be traced. Nitrous acid cannot be absorbed by soda-lye without loss, as it is partially resolved into nitric acid and nitric oxide. The violet colouration which appears during the action of reducing agents upon a nitrose is due to the solution of nitric oxide in the acid, and is possibly the result of a very unstable compound of nitrogen and oxygen, between NO and N_2O_3 .

Organic Thio Compounds.—O. Wallach and H. Bleibtreu.—The authors have obtained a series of homologues of a compound which they recently prepared by the action of iodethyl upon the sodium compound of thiocetanilid.

Thiamids of the Oxalic Acid Series.—O. Wallach and P. Pirath.—Not suitable for abstraction.

Certain New Salts of Aniline.—Miles Beamer and F. W. Clarke.—A description of the chlorate, perchlorate, iodate, hydrofluorate, phthalate, and mono-, di-, and trichloracetate.

Remarks on Lithium Picrate.—M. Beamer and F. W. Clarke.—The authors obtain this salt by dissolving lithium carbonate in an alcoholic solution of picric acid.

Bilic Acid, a New Oxidation-product of Cholic Acid.—F. Egger.—The author obtains this substance by acting upon 30 grms. cholic acid with 60 grms. potassium bichromate with 32.5 c.c. concentrated sulphuric acid diluted with 8 vols. of water. Action is set up on the application of heat. The composition of the pure acid is $C_{16}H_{22}O_6$.

Derivatives of Oxy-propyl-benzoic Acid.—R. Meyer.—The principal of these derivatives is para-acetyl-benzoic acid.

H. Köhler's Observations on Mercuric Iodide.—K. Kraut.—The writer is not willing to accept Köhler's results on the melting-point of mercuric iodide.

Synthesis of Chrysen.—C. Graebe and H. Bungener.—The authors obtain chrysen by the same method which led to the transformation of stilben and dibenzyl into phenanthren.

New Synthesis of Desoxybenzoin.—C. Graebe and H. Bungener.—Chloride of phenyl-acetic acid was treated with an excess of benzol and with aluminium chloride.

Derivatives of Mucic Acid.—E. Seelig.—An extensive memoir on dehydro-mucic acid and its salts.

Sulphuretted Acids formed by the Introduction of Brom-benzol and Chlor-benzol into the Organism.—M. Jaffé.—Not suitable for abstraction.

Constitution of Indigo.—E. Baumann and F. Tiemann.—An essay on the constitution of sulphindoxyl acid, on the conversion of indoxyl into indigo, and on the constitution of the latter.

Oxidation-products of Quinin.—Z. H. Skraup.—Among these are quitenin, $C_{19}H_{22}N_2O_4 + 4H_2O$, formic acid, and quinic acid, $C_{11}H_9NO_3$.

Constitution of the Quinin Bases.—Z. H. Skraup.—A theoretical memoir, not susceptible of useful abstraction.

Synthesis of Iso-succinic Acid.—H. Züblin.—Malonic ether in contact with sodium gives off hydrogen in abundance, and when treated according to the usual method for preparing sodium-acetic ether, it gives a product which was converted into iso-succinic ether by the action of iodomethyl.

Determination of the Vapour-densities of Certain Inorganic Bodies at Elevated Temperatures.—V. Meyer and C. Meyer.

Lecture Experiments.—A. W. Hofmann.—This paper also requires the accompanying illustrations.

Action of Phosphor-penta-chloride upon the Oils of Mustard and Cognate Bodies.—A. W. Hofmann.—An account of the choro-phenyl-oxyphenyl and amido oils of mustard.

Action of Hydrochloric Acid (Dry) upon Terpens.—W. A. Tilden.—Terpens of the two classes boiling respectively at 156° and 176° , and which are widely different in their optical behaviour, can be thus converted into one and the same inactive hydrocarbon.

No. 10, 1879.

Formation of Cinchomeronic Acid from Quinin, and its Identity with Pyridin-dicarbonic Acid.—H. Weidel and M. v. Schmidt.—The authors have obtained and examined the acid and neutral soda salts and the neutral lime salt of this acid. Among the products of the oxidation of cinchonin with nitric acid formed simultaneously are cinchonic (carbo-quinolinic acid), quinolic acid, cinchomeronic and oxy-cinchomeronic acids.

Nitro-phenanthren and its Derivatives.—G. A. Schmidt.—The author took for his point of departure mono-nitro-phenanthren, and finds that this compound exists in three isomeric forms. Among the derivatives obtained are amido-phenanthren, α -mono-nitro-phenanthren-quinon, β -mono-nitro-phenanthren, and γ -mono-nitro-phenanthren.

Reply to a Reclamation of Priority concerning the Reaction of Metallic Chlorides upon Aromatic Hydrocarbons.—B. Aronheim.—A reply to Mr. Watson Smith.

A Correction.—J. Nessler.—A reply to Dr. Lunge's note on gypsiferous wines.

Poly-substituted Ureas.—W. Michler and C. Escherich.—This memoir (a continuation) appears as two distinct papers in which the authors describe the reactions of chlorocarbonic oxide and dimethyl-amin, of dimethyl-urea-chloride with dimethyl-amin, of chloro-carbonic oxide and mono-methyl-amin, and of ammonia upon methyl-phenyl-urea-chloride.

Notice on the Nitrising of Benzol-sulphanilid.—W. Michler and G. Blattner.—A preliminary notice of experiments, undertaken in order to obtain highly nitrised nitro compounds by treating sulphanilides with fuming nitric acid.

Behaviour of Amines with Sulpho-chlorides.—W. Michler and G. Moro.—In this, the first section of their treatise, the authors describe the reaction of trichlor-methyl-sulpho-chloride and dimethyl-aniline. The resulting compound has a formula suitable for a tetra-methyl-diamido-benzo-phenon.

Phosphates of Zinc.—W. Demel.—The author has examined the acid zinc salts of phosphoric acid, and gives an elaborate crystallographical account of—
 $\text{ZnO}, \text{P}_2\text{O}_5, 2\text{H}_2\text{O} + 2\text{H}_2\text{O}.$

Analysis of a Turkey-red Oil.—G. Stein.—The author's method is to weigh 10 grms. of the sample into a porcelain capsule holding about 125 c.c., adding 75 c.c. cold saturated solution of common salt, and 25 grms. of dry wax. The whole is then heated on a water-bath. As turkey-red oil is insoluble in brine it rises in an anhydrous condition to the surface and combines with the melted wax. The cake of wax, when cold, is freed from brine by means of filter-paper, dried over sulphuric acid and weighed. On deducting the known weight of the wax the residue is the real oil.

Constitution of Sulpho-toluid.—R. Otto.—The author describes di-para-toluol-sulphide and di-para-toluol-sulphon.

Formulae of Quercitrin and Quercetin.—C. Liebermann and S. Hamburger.—The authors assign to quercitrin the formula $\text{C}_{36}\text{H}_{38}\text{O}_{20}$, which agrees well with that

of Hlasiwetz and of Rigaud. To quercetin they give $\text{C}_{24}\text{H}_{16}\text{O}_{11}$, the analysis likewise agreeing closely with those of Rigaud, Hlasiwetz, and Zwenger and Droncke. They have also formed and examined a number of the derivatives of quercetin.

Mercuric Chloro-iodide.—H. Köhler.—The author has established the real existence of this compound, HgClI , as a chemical individual.

Constitution of Indigo.—E. Baumann and F. Tie-mann.—The authors consider it necessary to ascertain whether indigo is a derivative of diphenyl or of diatyrenyl, and whether it contains imid groups.

Determination of the Vapour-density of Certain Chlorides.—V. and C. Meyer.—The vapour-densities here ascertained are those of stannous chloride, zinc chloride, and ferric chloride. Concerning the first of these compounds they consider it established that it should be represented not by the more simple formula, but by that corresponding to the molecular magnitude Sn_2Cl_4 .

Conversion of Furfuralgellic Acid into Azelalnic Acid.—P. Tonnies.—The ultimate step in this transformation is the reduction of butyro-furonic acid to azelalnic acid.

Relations of Dibrom-pyro-mucic Acid to Mucobromic Acid.—P. Tonnies.—A hypothetical paper.

Bulletin de la Société Chimique de Paris,
No. 3, 1879.

New Method of Determining Sulphur in certain Native Sulphides.—Albert Colson.—The author proposes to burn pyrites in a current of oxygen, and to titrate the products of combustion. The operation is carried on in a tube of green glass, placed in a furnace for organic analysis. One end of the tube is sealed, and to the other is fitted a stopper with two holes. One of these serves for the escape of the gaseous products of combustion which are received in a Liebig's bulb apparatus containing standard caustic soda. Through the other hole passes a long narrow tube, which conveys oxygen, free from water and carbonic acid, to a small platinum boat placed near the closed end of the combustion-tube, and containing half a gramme of the pyrites spread out in a thin layer. A plug of asbestos is placed about the middle of the tube, to avoid projections. Heat is first applied towards the open end of the tube, and as it approaches redness it is gradually extended towards the closed end. The current of oxygen is regulated so as to be always in excess. The disappearance of the white vapours formed in the bulb-tube during the operation is a sign that the sulphurous acid is expelled from the combustion-tube. The open end of the tube and the stopper are then washed, and the washings are added to the liquid in the bulb-tube, the sulphurous acid in which is then determined in the usual manner. If the pyrites contain carbonates, standard solutions cannot be used. It is then necessary to oxidise the sulphurous acid and determine the sulphuric acid formed.

Thermic Formation of Hydrogen Silicide.—J. Ogier.—The union of $\text{Si} + \text{H}_2$ is accompanied with a disengagement of heat to the extent of $+24.8$ cal.

Thermic Formation of Silicic Ether.—J. Ogier.—The heat of formation of silicic ether from pure alcohol and silica dissolved in water is 11.44 cal.

A New Derivative of the Parabanic Series.—E. Grimaux.—The substance in question is the amide of an oxalyl-biuretic acid, comparable to oxaluric acid.

Neutral Sugar and Inverted Sugar.—P. Horsin Déon.—Neutral sugar and inverted sugar have the same composition, equal parts of glucose and levulose. Neutral sugar is an inverted sugar in which the glucose possesses its maximum rotatory power. Its formation is the first step towards the inversion of sugar.

Palm-Sugar from Calcutta.—P. Horsin Déon.—This sugar is composed chiefly of cane-sugar, along with reducing sugar, gum, and mannite.

Researches on the Bacillus Ferment of Urea.—P. Miquel.—*Bacillus ureæ* belongs to the class of beings which M. Pasteur names *Anærobia*. At the end of its life it is resolved into brilliant, slightly elliptical, spores, which can resist for several hours a moist heat of 95° to 96°.

Hydrosulphuric Fermentation.—P. Miquel.—The author considers that there exists at least one organism capable of hydrogenising sulphur to an appreciable extent. This *Microbium*, which is met with in abundance in sewage, in drinking-waters, and even sometimes in rain-water, attacks insoluble albumen, and eliminates the bulk of its sulphur in the state of free hydrogen sulphide.

La Correspondance Scientifique.
No 63, October 7, 1879.

This issue contains neither chemical nor physiological matter.

MISCELLANEOUS.

City and Guilds Institute.—The City and Guilds of London Institute for the Advancement of Technical Education, announce the opening of their Technical Classes, at Cowper Street School, Finsbury. In the Section of Applied Physics, Mr. W. E. Ayrton will deliver a course of twelve lectures on "Some of the Practical Applications of Electricity and Magnetism," commencing Monday, November 3rd, at 7 p.m. In that of Applied Chemistry, Dr. H. Armstrong, F.R.S., will deliver a similar course on "The First Principles of Chemistry," commencing Wednesday, November 5th, at 8 p.m. An Inaugural Lecture will be delivered by Mr. Ayrton, on Saturday, November 1, at 8 p.m., on "The Improvement Science can Effect in our Trades, and in the Condition of our Workmen." A syllabus of each Course of Lectures can be obtained at the Halls of the Companies of Mercers, Clothworkers, and Drapers, or at the Schools, Cowper Street, Finsbury. Intending students should send in their names to the Demonstrator, Cowper Street Schools, Finsbury, E.C. The Inaugural Lecture will be free.

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THE CHEMICAL NEWS.

VOL. XL No. 1040

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SEPARATION AND DETERMINATION
OF MANGANESE.

By Professor J. VOLHARD.

The following account of this process is condensed from *Liebig's Annalen*. The author makes use of a standard solution of potassium permanganate, which he prepares by dissolving 38.5 grms. crystals of permanganate in about 2 litres water in a boiling-flask. The solution is poured into a well-stoppered bottle which holds 10 litres, filled up to the mark with water and well mixed by shaking. This store is kept in a dark place. To fill the burettes, the author uses a 1-litre washing bottle with a wide mouth covered with black paper. Before filling this vessel out of the stock-bottle, it is several times washed out with a little of the solution. The value of the permanganate solution is then determined, which is best effected iodometrically. For this purpose there are required a solution of sodium hyposulphite and a starch solution, both of known strength, besides a solution of potassium iodide and one of starch.

The reducing liquid is obtained by dissolving 30.061 grms. pure crystalline sodium hyposulphite along with about 3 grms. ammonium carbonate and making up to 1000 c.c.; 1 c.c. of this solution = 2 m.grms. manganese. To prepare the iodine solution, 1.5394 grms. dry iodine in crystals are dissolved in a little water by the aid of 3.5 potassium iodide and made up to 1000 c.c., 10 c.c. of this solution are reduced by 1 c.c. of the hyposulphite solution. The potassium iodide solution should contain per litre 55 grms. potassium iodide free from iodate. The solution of starch is prepared fresh daily.

About 10 c.c. of potassium solution are placed in a beaker, 4 to 5 c.c. pure hydrochloric acid with 150 to 200 c.c. water are added, and constantly stirring about 20 c.c. of the permanganate solution are added from a Gay-Lussac burette. The beaker is then placed under the Mohr's burette containing the hyposulphite solution, and some of this is added till the brown colour of the iodide solution is changed to a faint yellow. Starch solution is now added, and hyposulphite again till the blue colour is completely destroyed. In order to determine the small excess of hyposulphite which has been added, solution of iodine is added with a small graduated pipette till the blue colour just reappears. The volume of hyposulphite consumed, less one-tenth of the iodine solution used and divided by the volume of the permanganate solution gives the factor of the permanganate: 1 c.c. permanganate indicates 2 m.grms. $\times$ factor for manganese.

The standard of the hyposulphite solution should be checked from time to time. For this purpose the author uses a liquid prepared by dissolving 5.963 grms. potassium dichromate, and making up to 1000 c.c. Of this solution 1 c.c. gives up to hydriodic acid exactly as much oxygen as the quantity of permanganate, which indicates 2 m.grms. manganese; 20 c.c. of the chromate solution are measured off with a pipette, and let flow into the acidified solution of potassium iodide, proceeding exactly as directed above for standardising the permanganate. If more than 20 c.c. of the hyposulphite are used for reducing the iodine set free the hyposulphite was impure.

Any other permanganate solution of known strength will serve as well for the titration of manganese, and if a solution is kept in stock for the titration of iron, it would be superfluous to prepare another. It must be remembered, however, that only three-fifths of the oxygen in the permanganate indicated by ferrous oxide, oxalic acid, or hydriodic acid comes into play in the oxidation of man-

ganous oxide. The manganese standard may be found by multiplying the iron standard by 0.2946.

For the titration of manganese, if iron is absent, or present only in a small quantity, the solution of the salt of manganese is rinsed into a long-necked boiling-flask, with the addition of 1 grm. zinc sulphate, and diluted so that 100 c.c. may contain not more than about $\frac{1}{2}$ grm. manganese. If the solution is neutral, 2 or 3 drops of pure nitric acid are added (specific gravity 1.2); if acid, it is first neutralised with absolutely pure sodium carbonate, till a permanent precipitate begins to appear, then acidified with 3 or 4 drops of nitric acid and heated to a boil. The flask is taken from the fire and the permanganate run in with the burette, promoting the collection of the precipitate by vigorous shaking. The redness which appears towards the end, repeatedly disappears again on further agitation. If fine reddish-brown flocculi remain suspended in the liquid, so that it does not become thoroughly clear and transparent, the flask is set for a few minutes on a very gentle fire, but not allowed to boil.

The flocks become more compact and subside on shaking. The liquid at the end of the operation must have a distinct rose colour,* which must remain after repeated shaking. The zinc sulphate must be carefully tested; its dilute solution should not, on boiling, destroy the colour of a drop of permanganate.

Metallic alloys, such as cast iron and steel are dissolved for the determination of the manganese in dilute sulphuric acid with the addition of nitric acid. The solution is effected in a litre flask, heated on the water-bath. In a mixture of 3 volumes dilute sulphuric acid (1.13 specific gravity) and 1 volume nitric acid (1.4 specific gravity) common bottle-wire, on heating in the water-bath, dissolves in five or six minutes without violent evolution of gas. The digestion must be prolonged in order to convert the iron completely into oxide. Many ores and slags can only be dissolved by the aid of hydrochloric acid, in which case, after the oxidation of the iron, the hydrochloric solution is mixed with concentrated sulphuric acid and evaporated in a porcelain capsule, first in the water-bath and then in the gas-stove, till the sulphuric acid begins to escape. The mass is then rinsed with water into the litre flask. The more highly carburetted cast-iron, spiegel-eisen, and ferromanganese are dissolved in nitric acid in a small flask; the solution, without filtration, is rinsed into a porcelain capsule, evaporated to dryness in the water-bath, and heated in the gas-stove to the complete decomposition of the nitrates, whereby the carbonaceous matters are burnt. The residual oxides are digested with hydrochloric acid in a covered capsule on the water-bath, when the oxides dissolve. The hydrochloric acid is then again expelled by heating with sulphuric acid.

The bulk of the acid is next neutralised with pure sodium carbonate or hydroxide. Zinc white suspended in water is then added till all the iron is thrown down, which is ascertained by the circumstance that the solution suddenly coagulates and the supernatant fluid becomes milky. The flask is then filled up to the mark with water, allowed to settle for some minutes, and filtered through a dry folded filter into a dry vessel. A part of the filtrate (200 c.c.) is placed in a boiling-flask, acidified with 2 to 4 drops nitric acid and heated to a boil. The flask is then taken from the fire, and the permanganate run in with the burette. The titration can be repeated with a second and a third portion of the liquid.—*Liebig's Annalen der Chemie*.

Technical Education.—A class in connection with the City and Guilds of London Institute for the Advancement of Technical Education, for the study of blowpipe analysis and assaying, will be commenced on Wednesday next at the Birkbeck Institution, by Mr. G. Chaloner, F.C.S.

* The Germans understand, under "rosa," a colour rather inclining to the violet side of red.

ON THE CHEMICAL COMPOSITION OF
AMBLYGONITE.*

By SAMUEL L. PENFIELD.

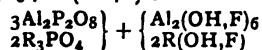
THE new mineral species triploidite described by Messrs. Brush and Dana† is shown by them to be isomorphous with wagnerite and closely related in composition to tripelite. These three minerals have respectively the formulæ $(\text{Mn,Fe})_3\text{P}_2\text{O}_8 + (\text{Mn,Fe})(\text{OH})_2$, $\text{Mg}_3\text{P}_2\text{O}_8 + \text{MgF}_2$, and $(\text{Fe,Mn})_3\text{P}_2\text{O}_8 + (\text{Fe,Mn})\text{F}_2$. From a comparison of these formulæ it is argued (*l. c.*, p. 45) that the relation between the minerals requires the assumption that the hydroxyl in triploidite must play the same part as the fluorine in the other two.

In this paper the author wishes to show that in amblygonite the hydroxyl group is also isomorphous with fluorine, and that in chemical composition the original amblygonite does not differ from the American and Montebbras varieties, which have been called hebronite. The results of his analyses require the adoption of a new formula for the mineral more simple than that previously accepted. For analysis he selected specimens from the three localities in Maine, from Branchville, Connecticut, where the mineral has been lately discovered by Messrs. Brush and Dana, also two varieties from Montebbras and one from Penig, Saxony, from a specimen in the Yale College collection.

The ratios from the analyses are collected in the following table by themselves, where R equals Li and Na:—

	P	Al	R	(OH,F)
I. Penig, Saxony . . .	1.00	0.96	0.98	1.16
II. Montebbras, France, A . . .	1.00	0.97	0.98	1.17
III. Auburn, Maine . . .	1.00	0.96	0.97	1.06
IV. Hebron, Maine, A . . .	1.00	0.97	0.95	1.13
V. Paris, Maine . . .	1.00	0.96	0.97	1.17
VI. Hebron, Maine, B . . .	1.00	0.98	0.95	1.27
VII. Branchville, Conn. . .	1.00	0.97	0.96	1.09
VIII. Montebbras, France, B . . .	1.00	0.96	0.96	1.21

It will be seen that all of these approach closely to the ratio 1 : 1 : 1 : 1, hence the author proposes the formula $\text{Al}_2\text{P}_2\text{O}_8 + 2\text{R}(\text{OH,F})$ or—



as the true formula for all varieties of this mineral.

Des Cloizeaux, from a difference in optical characters made out by him, has divided the mineral into two species: the original amblygonite, including I. and II. in the above list; and a second species for which he proposed the name *montebbrasite* (hebronite of von Kobell), including analyses III. to VIII. The mineral from Branchville has not been examined optically and the material is very unfavourable for such an examination. Owing to the close identity in chemical composition, it seems that a slight variation in optical properties is hardly sufficient ground for dividing the mineral into two species; but, on the contrary, the author thinks that the old name amblygonite should be retained, and that all varieties should be included by it.

It will be seen, in comparing the above ratios, that in every case the ratio of P to (OH,F) is in excess of that of the Al to R. Two theories suggest themselves to account for this, the first of which seems the most plausible. First, most minerals which are ordinarily regarded as anhydrous contain a small amount of water, which is not calculated in the ratios, and which is not regarded as essential to the composition. Now, if these minerals contain a small quantity of such accidental water, it will bring up the ratios very considerably, owing to the small molecular weight of water, and if the slight variation between

P, Al, and R be regarded as due to error of analysis, the excess of (OH,F) would be easily accounted for. Second, if we regard the difference between P, Al, and R as not due to error of analysis, and the fact that the variation in all is so constant suggests this, and regard enough of the water basic so that when added to the Al and R it will make the ratio with P equal 1 : 1 : 1, then the ratio of (OH,F) will be:—Penig, 1.02; Montebbras, A, 1.06; Auburn, 0.91; Hebron, A, 0.99; Paris, 1.02; Hebron, B, 1.18; Branchville, 0.97; Montebbras, B, 1.05. This relation seems rather striking, and although it is not as simple as we should like to have it, or perhaps as plausible as the first theory, yet it may possibly be the correct one. Whichever of these explanations is accepted, it will not materially alter the formula above made out for the minerals, the variation from which is too slight and not constant enough to be expressed by any different formula. The analyses show that water is found in the Penig and Montebbras varieties which have been regarded as anhydrous by some analysts. This may have been overlooked, and it is worth noting that in Plattner's "Blowpipe Analysis" the statement is made, of the Penig mineral, that water is expelled by heating in a closed tube. It will also be seen that these analyses differ from the older ones in that they are lower in alumina and higher in alkalis.

Method of Analysis.

Water was determined by ignition with oxide of lead in a porcelain crucible; it is completely driven off only by strong ignition, and it was found necessary to fuse the contents of the crucible over the blast lamp before constant results could be obtained. Fluorine was determined by decomposing a mixture of the mineral and powdered quartz with sulphuric acid, converting the silicon fluoride formed into hydrofluosilicic acid, precipitating the hydrofluosilicic acid with potassium chloride, and titrating the liberated hydrochloric acid with a standard alkali solution.* The varieties from Penig and Montebbras are decomposed only by prolonged action of sulphuric acid. Phosphoric acid was determined by fusing the mineral with sodium carbonate, boiling out the fused mass with water and dilute nitric acid, nearly neutralising the excess of acid with ammonia, precipitating with molybdic solution, and then proceeding in the usual way.

To determine the bases, one gramme was weighed into a large platinum crucible, mixed into a paste with from two to three c.c. of sulphuric acid, and heated, with the crucible covered, over a low gas flame till not over a c.c. of sulphuric acid remained. The contents of the crucible were then rinsed into a platinum evaporating dish, and treated with a quantity of strong hydrochloric acid; after heating and concentrating a clear solution was obtained. The excess of acid being removed by evaporation, the contents of the dish were rinsed into a beaker, filtered where necessary, the undissolved portion after incinerating the filter paper was treated with hydrofluoric acid, then with a drop of sulphuric acid; the hydrofluoric acid expelled by evaporation and the solution added to the other solution of the bases. To obtain the bases as chlorides the sulphuric acid was precipitated from the solution with barium chloride, and the barium sulphate filtered off. The solution was then heated to boiling, and a hot solution of barium hydroxide added; this precipitated all the phosphoric acid and part of the alumina. The solution contained all the lithia and most of the alumina which went into solution in the excess of barium hydroxide. After filtering and washing the precipitate it was dissolved in hydrochloric acid, the excess expelled by evaporation; the residue taken up in a few drops of hydrochloric acid and water, and poured when hot into a boiling solution of sodium hydroxide and a little barium hydroxide in a platinum dish; this again precipitated all the phosphoric acid, while the alumina went into solution in the alkaline hydroxides. After filtering, the filtrate was acidified with

* Abstract of a paper in the *American Journal of Science*,
† *Am. Jour. Sci.*, III., xvi., 42, July, 1878.

* See Remsen's *Am. Chem. Jour.*, vol. i, No. 1.

hydrochloric acid, the barium precipitated with sulphuric acid, and the alumina precipitated with ammonia. The alumina found at this point amounted usually to about one-half per cent. The insoluble barium phosphate containing also traces of iron, manganese, and calcium was dissolved in hydrochloric acid, the barium precipitated with sulphuric acid, filtered, and the filtrate made alkaline with ammonia; this precipitated any iron, manganese, or calcium as phosphate, which was filtered off and examined separately. It was intended at this point to determine the phosphoric acid by direct precipitation with magnesia mixture, but the results coming out too low an examination of all the barium sulphate precipitates showed that a quantity of the phosphoric acid had been precipitated along with the barium sulphate. The first filtrate from the barium hydroxide precipitate contained, free from phosphoric acid, all the lithia, the larger part of the alumina, and the excess of barium. It was heated to boiling and ammonium carbonate added; this precipitated the barium and aluminium, and left the lithia in solution. The precipitate was washed first by decantation, then with hot water on the filter pump; it was not considered free from lithia, however, as it is practically impossible to wash a large barium carbonate precipitate free from lithia. The filtrate was evaporated to dryness, the ammonia salts expelled by ignition, traces of barium separated a second or third time when necessary, evaporated to dryness again, lithia separated from soda and potash by means of absolute alcohol and ether, the lithia weighed as sulphate, and the soda and potash as chlorides. The chlorides were tested carefully for potash by evaporating with excess of platinum chloride and taking up in alcohol. The precipitate produced by ammonium carbonate was dissolved in hydrochloric acid, the barium precipitated with sulphuric acid, filtered, and alumina precipitated in the filtrate with ammonia, the precipitate was washed with hot water, ignited finally over the blast lamp to expel sulphuric acid, and weighed as oxide. The filtrate from the alumina was regarded as containing a trace of lithia which had been retained by the barium carbonate precipitate, it was evaporated to dryness, the ammonia salts expelled by ignition, taken up in water, filtered into a weighed crucible, evaporated to dryness and weighed, the lithia found amounted to from one-quarter to one per cent.

The solutions were kept as far as possible from all contact with glass, the evaporations being carried on in large platinum dishes. The reagents were carefully selected and purified. Sodium hydroxide free from aluminium and silica was obtained, prepared from metallic sodium. Owing to the limited amount of material from Penig only three-quarters of a gramme was used in the determinations, and duplicates of the water and fluorine determination were not obtained. For the occurrence and associations of amblygonite at Branchville, Connecticut, see the papers by Messrs. Brush and Dana.*

Sheffield Laboratory, June 18, 1879.

ANALYSIS OF CINCHONA BARKS.

A CORRESPONDENT favours us with the following valuable information on this subject, in answer to a question which appeared lately in our "Notes and Queries" column.

"ETHER PROCESS,"

FOR THE ESTIMATION OF QUININE IN CINCHONA BARK.

Take 1000 grains finely-powdered bark; 8 fluid ozs. of alcohol (methylated) at 35 to 40 per cent, sp. gr. 0.872; 500 grains hydrate of lime; 16 fluid ozs. ether (methylated), 0.730 sp. gr.

Let the bark be powdered without leaving a residue, and passed through a sieve of at least 100 meshes to the

* *Am. Jour. Sci.*, July and August, 1878, and May, 1879.

inch (fine cypress). Pour upon the powder sufficient alcohol to form a liquid paste, heat for a few minutes until the fibre is thoroughly penetrated by the liquid. Introduce into the paste the hydrate of lime in fine powder, mix thoroughly, heat on a plate until all the alcohol is expelled and the powder thoroughly dry. Treat this in a displacement-apparatus with successive portions of ether; thus 4 ozs. to macerate, and six successive 2 ozs.; evaporate percolate rapidly.

1. Fuse product at 120° C. = crude quina + colouring- and fatty-matter.

2. Dissolve the residue in alcohol, and estimate with standard sulphuric acid = total sulphate of crystalline and amorphous quina.

3. Weigh the sulphate of quinine crystals obtained.

Cinchona barks of all species contain a natural acid, called by some kinic and by others quinovic acid. This acid exists in combination with one or more of the alkaloids, quinia, quinidia, cinchonina, cinchonidine, according to the variety of the bark.

Calisaya Bark (Bolivia), of both quilted and flat varieties, contains wholly, or almost wholly, quinia; quinidia and cinchonina being rarely met with. (Ether process specially adapted.)

Red Bark, Cartagena, Soft Columbian, by which names they are known in commerce, contain both alkaloids of quinia and cinchonina, and to a greater or less degree quinidia. The ether process is also applicable, in so far as the estimation of the quinia and consequent determination of the commercial value for the manufacture of quinine is concerned.

Loxa, Brown, or Grey Barks, commercial names, contain almost wholly cinchonina and cinchonidine. The ether analytical process is not applicable to this variety.

The lime is added to decompose these quinovates of the alkaloids, in order that the latter may be exposed to the solvent action of the menstruum in the free state.

The ethereal solution (percolate) will contain the total quinia with traces only of the other alkaloids when they are present, together with fatty- and colouring-matter.

Evaporated ethereal solution, fused at 125° C., will be the total crude quinia plus colouring- and fatty-matter. Note the weight.

Dissolve by aid of a water-bath in about 1 oz. of absolute alcohol, and add gradually (having a piece of litmus paper in the solution), until a faint acid reaction is indicated, a standard sulphuric acid of known strength, by preference one in which 100 c.c. = 10 grms. crystal sulphate quinine, if a 1000 grain measure burette be used, so that 1000 = 100 grains quinia sulphate crystals.

Note quantity of acid for neutralisation.

Product estimated as total basic sulphates of crystalline and amorphous quinia. The strength of the acid can easily be deduced from the following:—

Composition of air-dried crystal sulphate of quinia—

74.31 anhydrous quinia.	
(SO ₃ —) 9.17	11.23 mono-hydrated acid.
2.06	
14.45 water lost at 100° C.	

100.00

If pure sulphuric acid of 1.843 sp. gr. = 96.8 per cent of H₂SO₄ real, then 11.23 = 11.6 acid of sp. gr. 1.843.

Take, therefore, 11.6 grms. and dilute to a litre with water

To this alcoholic basic sulphate, when evaporated to dryness, add as many measures of the standard acid as used in the first instance. This will convert basic sulphate of quinine into acid sulphate, a salt exceedingly soluble (unlike ordinary sulphate, which will crystallise in the filter). Add about 1 oz. of water, and boil with a spirit-lamp, being careful that no portion on the sides of the dish (No. 1.) become charred.

Solution of the salt being complete add a small quantity of animal charcoal (about 5 per cent of the bark originally

used). This animal charcoal should be completely purified by previous digestion in hydrochloric acid until the phosphate of lime is nearly wholly removed. Digest for about ten minutes, or longer if time is no object, then filter into a small dish (No. 11.), wash the filter with two successive portions of boiling water (boiled in dish No. 1.) acidulated with about a drachm (to cleanse it of all alkaloid) of the standard sulphuric acid.

The filtrate will consist of a decolourised solution of acid sulphate of quinine, strongly acid to litmus, and on the filter will be left with the animal charcoal all colouring- and fatty- (tarry?) matter.

Now evaporate the liquors (on a water-bath if time will permit) to about 1 fluid oz., or more if the bark is very rich in alkaloid (such as in quilted Calisaya barks for instance); raise to boiling-point, and add *very cautiously*, drop by drop, solution of ammonia of about 3 per cent (1 part liq. ammon. 0.880, 10 parts water), until the solution has but a faint acid reaction, at which stage the quinia salt will again be as a basic sulphate; set aside to cool. When cold the whole will be a mass of crystals. Throw upon a dry filter, and allow the mother-liquor to drain away. This liquor will contain the sulphate of amorphous quinia, any traces of the sulphates that may have been present, of cinchonina, cinchonidine, and quinidia, with sulphate of ammonia.

If the experiment has been properly conducted no crystalline sulphate of quinine will be in this mother-liquor, as *sulphate of ammonia* decreases the solubility of sulphate of quinine.

Until, however, the operator becomes perfectly familiar with the process it would be advisable to evaporate this mother-liquor, neutralise any excessive acidity, and look for a further crop of crystals, which will rarely exceed 0.1 per cent of the bark operated on.

Dry crystals in air; weight = amount of crystalline sulphate of quinine in original bark. A more rapid and certain method is to dry them at 100° C., and estimate as 85.5 dry = 100 crystals (vide composition of sulphate of quinine).

Any method of analysis intended as a sure guide for the manufacture of quinine must embrace a stage in which the quina is weighed as *pure white sulphate*.

Advantages of this Process as Compared with Acid Process.

1. Rapidity (twenty-four hours at the most.)
2. Extraction of that alkaloid only upon which the commercial value of the cinchona barks depend.
3. Non-production of amorphous quinia, so liable by protracted boiling, as in acid process.
4. Inexpensive.

"ACID PROCESS,"

FOR THE COMPLETE ANALYSIS OF ANY VARIETY OF CINCHONA BARK.

Take of bark in fine powder 1000 grains; sulphuric acid (1.843), 50 grains; water, 2 pints; hydrate of lime, a sufficiency.

Add the acid to the water, and divide it into two portions, in which successively boil the bark. After each decoction throw upon a calico filter until drained; then boil the bark in $\frac{1}{2}$ pint of water only, drain upon filter, and press. Mix the decoctions, concentrate to $\frac{1}{2}$ pint, cool, and add milk of lime until the liquid is distinctly alkaline. Collect the precipitate by filtration, and purify by careful washing, then dry at 95° to 100° C., and powder. Boil this powder repeatedly in strong alcohol (about 90 per cent), and evaporate alcoholic extracts to dryness. Boil the residue with acidulated water, and filter. Make the solution alkaline with caustic soda (this alkali is to be preferred, as quinia is insoluble in soda but soluble slightly in potash or ammonia) in a separator, and shake with chloroform. Draw off chloroform solution, and evaporate in a tared capsule. Product = total alkaloids of quinia, cinchonina,

quinidia, &c. (All the alkaloids are soluble in chloroform.)

Treat the alkaloids with ether. Evaporate ethereal solution. Product = amorphous and crystallisable quinia. Treat this for obtaining crystalline value as directed in the ether process.

The insoluble portion is then dissolved in dilute acetic acid, and treated with iodide potassium (sat. sol.); the precipitate, if any, is iodide of quinidia: can be estimated thus:—

100 grains = 71.68 quinidia

= 94.5 quinidia sulph.

The liquid treated with ammonia or soda gives a precipitate of cinchonina, which, when dried at 120° C., may be estimated as—

82 = 100 cinchonina sulphate crystals.

Another Method, although less accurate, is to digest in proof-spirit (alcohol 0.920 sp. gr.), which dissolves quinidia, and leaves undissolved the cinchonina and cinchonidine.

If these latter alkaloids are required in a quantitative crystalline form dissolve the residue, after treatment with ether, in a mixture of one volume of alcohol 0.795 and 9 vols. of ether; from this the quinidia will crystallise. The cinchonina can afterwards be crystallised from a saturated alcoholic solution.

To this process might be appended another, by percolation in the cold with alcohol. But as this process occupies considerable time in its first stage, it cannot be recommended for commercial purposes.

EXAMINATION OF THE NORTH CAROLINA URANIUM MINERALS.

By F. A. GENTH.

PROF. W. C. KERR describes the occurrence of uranium in Mitchell County, North Carolina, as follows:—"A new locality, the Flat Rock mine, recently visited, has yielded the following in immediate association, viz., uraninite, gummite, uraconite, and, as incrustations on the outside of the latter, and of the fragments of rock adjacent, torbernite and autunite. These minerals occur only in one part of the mica-bearing portion of a very large granite vein, and are found in irregular nodules and rounded masses, some with a nucleus of uraninite of $\frac{1}{4}$ to $\frac{1}{2}$ inch, enveloped with a heavy layer of gummite, outside of which is a pale, yellow, earthy coating from $\frac{1}{4}$ to $\frac{1}{2}$ of an inch thick, which is uranochre or uraconite. One lump, the largest, weighs just a pound, and in all, I obtained between 3 and 4 pounds. The quantity of pitchblende remaining unaltered is very small, and by far the greater part of the mass of the nodules, probably nine-tenths, is gummite; and the smaller ones are nearly or entirely changed to uraconite."

Through the kindness of Prof. Kerr I have recently come into possession of specimens of this highly interesting occurrence, which fully agree with the description above quoted. As his mineralogical determinations, however, were not supported by chemical analysis, and, especially as the composition of gummite is very doubtful, I thought that an investigation of the uranium minerals from this locality would be desirable.

I regret that of the uraninite no material for analysis could be obtained. One of my specimens of gummite, however, still contains one or two small fragments of uraninite; the larger one has a brownish black colour on the margin, changing into brownish, and, by degrees, into the pure gummite. It is also penetrated by small veins, showing the gradual alteration of the uraninite.

Uranotil.—E. Boricky.

The analysis of the pale yellow coating surrounding the gummite shows that it is a variety of *uranotil*, and not

uranochroa or *uranocomite*. The original uranotil has been found in cavities of quartz, associated with fluorite, at Wölsendorf, in Bavaria, in lemon-yellow rhombic needles, of a specific gravity of 3.95. For comparison I will give below Boricky's analysis of it.

The North Carolina variety is apparently amorphous; massive, compact. $H = 2.5$. Sp. gr. = 3.834. Lustre, waxy to dull; colour, from pale straw-yellow to lemon-yellow; streak, pale straw-yellow; opaque; fracture, uneven.

B. B. in a tube yields water, and becomes brownish yellow; with fluxes the uranium reactions. Easily soluble in chlorhydric acid; on evaporation yields a jelly of silicic acid.

The analyses gave, after deducting a small quantity of quartz, mica, and felspar:—

			Mean.	Boricky.	Calculated.
SiO ₂	13.55	13.88	13.72	13.78	13.95
Al ₂ O ₃ , Fe ₂ O ₃	traces	traces	traces	0.51	—
UO ₃	66.76	66.59	66.67	66.75	66.98
PbO	0.74	0.45	0.60	—	—
BaO	0.28	0.48	0.28	—	—
SrO	0.13	—	0.13	—	—
CaO	6.23	7.11	6.67	5.27	6.51
P ₂ O ₅	not det'd	0.29	0.29	0.45	—
H ₂ O	not det'd	12.02	12.02	12.67	12.56
		100.82	100.38	99.43	100.00

Rammelsberg* gives for uranotil the formula—



My analyses agree far better with the following formula— $Ca_3(UO_2)_6Si_6O_{21} + 18H_2O$, as will be seen from the above calculated analysis. It forms an incrustation upon gummitite and results from its alteration, sometimes (as has been observed by Prof. Kerr) changing the entire mass of the latter, and yielding nodules of pure uranotil; mostly, however, converting not only the outside of the gummitite into uranotil, but penetrating its whole mass, so that even the purest particles contain already a large percentage of it, as will be seen from the analyses of the gummitite.

Gummitite.

The orange-coloured mineral from the Flat Rock mine has been called gummitite, from its resemblance to that from European localities. It occurs in amorphous, compact, nodular masses. $H = 3$. Sp. gr. = 4.840. Lustre, faintly resinous to dull; colour, reddish yellow to deep orange-red; streak, orange-yellow; opaque; fracture, subconchoidal to uneven.

B. B. yields water and turns reddish brown; upon charcoal with sodium carbonate yields metallic lead; with fluxes gives the reactions of uranium. Easily soluble in acetic acid.

Below I give the analyses of the purest deep orange-red material, after deducting a minute quantity of quartz, felspar, and mica. For comparison I add the only analysis which is known of gummitite, that from Johann-Georgenstadt in Saxony, by Kersten.

				Mean.	Johann-Georgenstadt.	Kersten.
SiO ₂	4.49	4.83	4.58	4.63	—	4.26
Al ₂ O ₃	0.67	0.40	—	0.53	Mn ₂ O ₃	0.05
BaO	0.88	—	—	—	As, Fl	traces
SrO	0.05	1.12	[4.59]	1.08	—	—
CaO	1.96	2.14	—	2.05	—	6.00
PbO	5.48	5.58	5.64	5.57	—	—
UO ₃	[75.71]	75.50	74.39	75.20	—	72.00
P ₂ O ₅	0.12	0.07	0.16	0.12	—	2.30
H ₂ O	not det'd	10.43	10.64	10.54	—	14.75
		100.07	99.72	99.36		

The opinion of chemists as to the composition of gummitite is very much divided. Kersten thought it to be a

* *Mineralchemie*, 692.

combination of phosphate of lime with uranic hydrate, the composition of which he expressed by the formula—



Berzelius considered it as a mixture of basic phosphate and basic silicate of uranic oxide and lime. Hausmann takes it to be principally the hydrate of uranic oxide. Rammelsberg expresses the same opinion, but believes that the analyses of gummitite, together with those of eliasite and pittinitite, represent mixtures, and that from their analyses no rational composition can be derived. Hermann, on the other hand, considers these and other uranium minerals, in which silicic acid has been found, even uraninite, as definite compounds of the formula— $4RO, SiO_2 + 4(4R_2O_3, SiO_2) + mH_2O + nX$. The accessory molecule X being wanting in pittinitite, and in the other species represented by $R(AsS)$ in uranochalcite, by $4RO, U_2O_3$ in uraninite, by CaO, CO_2 in eliasite, by $3CaO, P_2O_5$ in phosphor-gummitite, and by $3CaO, (PV)_2O_5$ in vanadin-gummitite. Hermann's views are as untenable as those of the others. Nearest to the truth, in my opinion, comes Paterson, who holds that silicic acid and phosphoric acid are not essential, and that gummitite is principally a lime-uranate, $CaO, 2U_2O_3 + 6H_2O$, analogous in composition to the artificial uran-yellow.

Gummitite is the result of an alteration of uraninite, and it seems to me that both pittinitite and eliasite are intermediate between the two, containing more or less of either. But I have already pointed out the fact that the gummitite from North Carolina is a mechanical mixture, and that uranotil penetrates the whole mass. The gradual change from gummitite into uranotil can be observed on every specimen from this locality.

If we calculate from the SiO₂ in the gummitite the requisite constituents of uranotil, making up the minute deficiency of CaO by its equivalent of BaO, we get:—

SiO ₂	4.63
CaO	2.05
BaO	0.30
H ₂ O	4.17

33.38 per cent of uranotil.

The PbO and BaO may be most rationally considered

as being present in the form of $M(UO_2)_2O_3 + 6H_2O$; we get therefore—

PbO	5.57
UO ₃	14.39
H ₂ O	2.70

22.66 per cent of lead-uranate; and

BaO	0.78
UO ₃	2.93
H ₂ O	0.55

4.26 per cent of barium-uranate.

There remain 35.65 per cent of UO₃, which is evidently present as uranic hydrate, requiring 4.45 per cent of water.

The gummitite from the Flat Rock mine is therefore a mechanical mixture of:—

Uranic hydrate, $H_2(UO_2)_2O_3 + H_2O$	40.10 p.c.
Uranotil, $Ca_3(UO_2)_6Si_6O_{21} + 18H_2O$	33.38 "
Lead-uranate, $Pb(UO_2)_2O_3 + 6H_2O$	22.66 "
Barium-uranate, $Ba(UO_2)_2O_3 + 6H_2O$	4.26 "
	100.40

The analyses of pittinitite and eliasite admit of no calculation, as they appear to have too many foreign substances present, and as the amount of uranic oxide which they evidently contain has not been determined; if we take the analysis of gummitite from Johann-Georgenstadt, we find it probably to be a mixture of calcium-uranate, with uranotil, a uranium-phosphate (phosphuranylite), and uranic hydrate. These calculations give the following amounts;—

S.O ₂		4.26
UO ₃		20.45
CaO		1.99
H ₂ O		3.84

30.54 per cent of uranotil.

CaO		4.01
UO ₃		41.25
H ₂ O		7.73

52.99 per cent of calcium-uranate.

P ₂ O ₅		2.30
UO ₃		4.68
H ₂ O		1.75

8.73 per cent of phosphuranylite.

UO ₃		5.62
H ₂ O		0.70

6.32 per cent of uranic hydrate.

The gummitte from Johann-Georgenstadt has therefore probably the following composition, corresponding with Kersten's analysis:—

Uranic hydrate, H ₂ (UO ₂)O ₂ + H ₂ O ..	6.32 p.c.
Uranotil, Ca ₃ (UO ₂) ₆ S ₆ O ₂₁ + 18H ₂ O ..	30.54 "
Phosphuranylite, (UO ₂) ₃ P ₂ O ₈ + 6H ₂ O ..	8.73 "
Calcium-uranate, Ca(UO ₂) ₂ O ₃ + 6H ₂ O ..	52.99 "

98.58

Phosphuranylite.—A new species.

Rhombic (?). Under the microscope very minute rectangular scales with pearly lustre can be distinguished. In pulverulent incrustations upon quartz, felspar, and mica. Colour, deep lemon-yellow.

B. B. in the tube yields water and becomes reddish brown while hot, brownish yellow after cooling; readily soluble in nitric acid, yielding with ammonium molybdate a yellow precipitate; contains no arsenic acid.

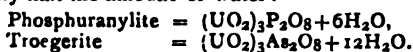
As this mineral is very rare, only a very minute quantity could be obtained which was free from autunite; this being in the form of a thin coating upon quartz, both together had to be taken as material for analysis.

0.6705 grm. contained 0.5617 grm. quartz—the analysis was therefore made with 0.1088 grm. of material. This was slightly mixed, probably with cerussite, which was visible under the microscope in the form of very small colourless particles. In the first column I give the results of the analysis, in the second the percentage of the phosphate, excluding the lead oxide, and in the third the calculated percentage, corresponding to the formula given below.

			Calculated.
UO ₃		71.73	76.71
PbO		4.40	—
P ₂ O ₅		11.30	12.08
H ₂ O		10.48	11.21

97.91

It will be seen from this analysis that the uranium and phosphorus are present exactly in the ratio of 3 : 2, and that the composition of phosphuranylite may be expressed by a formula analogous to that of troegerite, but containing only half the amount of water:—



Prof. Kerr mentions torbernite as one of the uranium minerals from the Flat Rock mine, and he sent me some uranite, which had a deep green colour and resembled it very closely; but the chemical examination of one of the darkest green crystals showed that it contained only lime, and not a trace of copper oxide, and that the uranite from this locality is therefore only autunite.—*American Chemical Journal*.

THE SIEMENS LIGHT AT THE
BRITISH MUSEUM.

ON Monday, the 20th inst., at 6 o'clock in the evening, a select party of officials, scientific men, and members of the press assembled in the Reading Room of the British Museum for the purpose of testing the new plan of lighting that building by the electric light as carried out by Messrs. Siemens, the electrical engineers. The main difference between the method adopted by Messrs. Siemens and that experimented upon by the Société Générale d'Electricité consists in the use of four very powerful lamps hung high over the heads of the readers, instead of a number of less powerful lights at a lower elevation. Two different systems of lighting are adopted, one for the Reading Room, and the other for the passages, hall, and court yard. The Reading Room is lighted by four lights of 5000 candle-power each, one being in the middle of the circle, and the other three arranged round it at equal distances in the form of an equilateral triangle. These four lights are produced by continuous currents passing through wires carried over the roof of the Reading Room and through the top of the glass lantern in the centre of the dome. The electricity is supplied by five dynamo-electric machines placed in a shed at the back of the Museum buildings at about 200 yards distance, and worked from two 8-horse-power semi-stationary engines, supplied by Messrs. Wallis and Stevens, of the North Hants Iron Works, Basingstoke.

The Reading Room regulators are the invention of Messrs. Siemens and Halske, of Berlin, and on Monday week they appeared to do their work thoroughly if we except a very unpleasant crepitation now and then, which a scientific man present aptly compared to the rale of a typhoid fever patient. There is also an occasional purple glow for a few seconds caused no doubt by some potassic impurity in the charcoal. These regulators are of a sufficiently simple kind and may easily be managed even by unexperienced hands. They each contain 19 inches of double carbon, which are consumed at the rate of 3 inches an hour. The lights consequently will burn for a little more than six hours without the regulators being interfered with. The other seven lights, of which two are placed in the Court Yard, one under the portico, one in the Hall, one in the passage leading to the Reading Room, one in the Sculpture Gallery, and the remaining one in the engine house, are worked by alternate current machines, and the regulators are actuated by two coils acting on a differential principle, one of them forming part of the main circuit and tending to separate the carbons, the other tending to bring them into contact. The position of the carbons depends therefore not upon the strength of the current, but upon the relative amount of electricity passing through each coil. These regulators are worked by two alternating current machines and are all in one circuit, the length of which is about 1200 yards, their construction being such that if one becomes extinguished either from accidental breakage or from want of carbon it does not affect the others. The carbons in these lamps burn for about five hours, and may be renewed in half a minute. The wires supplying the current for these regulators are laid partly in the basement of the building, and partly in pipes under the ground. The light given out by each of these lamps is equal to 600 candles, and their steadiness says much for the excellence of the principle upon which they are constructed.

The light from the lamps in the Reading Room passes through an opal glass lantern in the shape of an inverted truncated pyramid, a very small amount of the upward rays being thrown downwards by a circular reflector of opal glass. It is a singular fact that most gas and electrical engineers seem to take it for granted that this material is one of the best possible reflectors. For softening a too brilliant light it is unequalled, but its use as a reflector is almost nil. In the very powerful lamps erected by Messrs. Sugg in Waterloo Road and Waterloo Place,

the upper portion of the lantern consisted of a hollow pyramid of opal glass, which absorbed nearly the whole of the upward rays, while the small amount reflected downwards were thrown into the wrong place. In the present instance the whole of the light thrown upwards is lost with the exception of the very small proportion which is thrown down by the flat opal glass reflectors placed above the lamps.

The best method of distributing the electric, or indeed any very powerful light, over a given area has yet to be devised. The problem is a very simple one, but this is not the place for its solution. A proper knowledge of the most elementary laws of reflection will show that the plans at present in use are wrong in principle and involve a serious waste of lighting power.

The powers of the light were severely tested by several of the gentlemen present. Mr. George Bullen, the Keeper of the Printed Books, put it to several practical working tests in different parts of the room, and found that at the desks which are farthest from the centre there was a decided lack of light for those who, like Mr. Bullen, have somewhat impaired their sight by incessant literary labour, while at the catalogue desks and in the centre of the room, the light was much more than sufficient to read even the smallest print. He consequently made a most valuable suggestion to Dr. Siemens, which met with the immediate approval of that gentleman and of all present. The diameter of the Reading Room it will be remembered is exactly 140 feet, the half of which is 70 feet. As at present placed the three outer lamps hang at a distance of about 35 feet from the centre, and of course at an equal distance from the wall. A moment's consideration will show that according to this arrangement the central portion of the room gets an overplus of light, while the part near the wall is underlighted. Mr. Bullen consequently very properly suggested that the three outer lamps should be hung at a distance from the centre of two-thirds of the radius of the circle, by which means each third of the line between the centre and the circumference would be equally illuminated. By moving the outer lamps nearer the wall the outer row of the catalogues will be taken out of the deep shadow in which they were thrown by the arrangement adopted on Monday; it is to be hoped, therefore, that Mr. Bullen's suggestion will be carried into effect.

Dr. Carter Blake submitted the purity of the light to some very searching colour-tests, and found that the most delicate tints were as easily distinguishable as in bright daylight. This will be good news for those artistic workers who are so frequently rendered compulsorily idle during dark, or, worse still, yellow weather.

When the modifications indicated above are carried out there will be no possibility of doubting the thorough practicability of lighting the British Museum Reading Room by electricity, and readers of all classes will owe a heavy debt of gratitude to Mr. Bond for the persistency which he has shown in carrying out this vast improvement.

An important factor in the success of the experiment must be looked for in the engines supplied by Messrs. Wallis and Stevens, which work in an exceptionally steady manner, an advantage which is due to the use of an extremely sensitive form of governor, which keeps them at a uniform rate of speed, thereby securing almost perfect steadiness in the lights. Good as it is this form of governor has been further improved upon by Messrs. Wallis and Stevens. This new form of governor will be applied to the British Museum engines as soon as it has been thoroughly tested at Basingstoke. The peculiarity of the new governor lies not so much in its steadiness as in its allowing the engineer in charge to increase or diminish the normal speed of the engine at pleasure, according as the lights require it, and then to maintain with perfect regularity the particular speed which is found in practice to give the best light.

The best way of throwing down the upwards rays, most of which are now lost, should form the subject of serious

experiments. One thing is certain: opal glass is about the worst reflecting material that can be adopted. The dark colour of the interior of the Reading Room—the natural result of a long course of London fog and smoke—also causes a great waste of light. This is, however, a defect that can be very easily altered at a comparatively small expenditure of money and trouble.

In conclusion we must congratulate Mr. Bond on having carried his experiments to such a successful issue. In consequence of this success the Reading Room has been thrown open to readers every evening up to seven o'clock since Wednesday, the 22nd inst., and will continue open up to that hour until further notice. There is only one opinion amongst the readers as to the general efficiency and convenience of the light. It may be as well to mention that no books from the large library can be taken out after half-past four, but books already in the possession of readers before that hour may be kept until half-past six. The interior of the library not being lighted is the obvious reason for this regulation. The Library of Reference in the room itself is of course available up to closing time.

CORRESPONDENCE.

DISSOCIATION OF CHLORINE.

To the Editor of the Chemical News.

SIR,—A more careful reading of the few words from me under the above heading in the *CHEMICAL NEWS*, No. 1034, would make it perfectly clear that I did not confound the platinic and platinous chlorides, as suggested by Mr. Watson Smith in the *CHEMICAL NEWS*, No. 1035.

The presumption is that an investigator of such standing as Prof. V. Meyer, has taken all suitable precautions; yet in the case of so important a discovery as that in question, of which at the time of writing my only information was derived from the notice in *CHEMICAL NEWS*, No. 1027, any possible and less obvious source of error may well be brought to notice. And the question still remains, and is pertinent to such of V. Meyer's experiments as were made with platinous chloride; may not platinous as well as platinic chloride, the former presumably made from the latter, retain a portion of the compounds of oxygen and nitrogen at a temperature above that at which the lower chloride begins to be decomposed? assuming the original source to be as usual platinum and aqua regia, the solution afterwards repeatedly evaporated with hydrochloric acid and water. In the experiments of mine referred to, which were necessarily made under pressure, there was nothing to show that at such temperature oxygen compounds had ceased to be given off.—I am, &c.,

F. P. DUNNINGTON.

FLUID MEDIUM FOR IMMERSION LENSES.

To the Editor of the Chemical News.

SIR,—I find from conversation with those who have worked with the new "Homogeneous Immersion" (Oil Immersion) Lenses, that a better fluid medium is a desideratum. The essential oil used is too mobile, and consequently difficult to keep in its place between the front lens and the object, besides acting injuriously on some of the mounting varnishes.

Optically it answers well enough. The medium to be perfect should be of the same refractive and dispersive power as crown-glass,—thick enough to keep its place between the object-glass and the cover, not liable to evaporate rapidly or deposit crystals. Should not act upon glass, metals, or the usual cements employed in mounting objects.

As the introduction of these objectives—which certainly have great advantages optically—can hardly become general until a better medium is obtained, may I ask you to direct, as far as you are able, the attention of chemists to the subject? A suitable fluid may probably be easily attainable, but opticians and microscopists are probably the persons least likely to discover it.

The metals employed in mounting objectives are either brass or an alloy of nickel; platinum is not sufficiently rigid to bear turning to a thin edge.—I am, &c.,

W. T. SUFFOLK.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 15, October 13, 1879.

The Present State and the Future of Thermo-Chemistry.—M. Berthelot.—The author gives a history of thermo-chemistry and explains the plan of his new work, "Essay on Chemical Mechanics founded upon Thermo-Chemistry." The first volume of this work is devoted to calorimetry, that is to say, the measurement of the quantities of heat set in action in chemical processes. The second volume comprises the general study of chemical composition and decomposition, especially that of systems in equilibrium between two contrary tendencies. The remainder of the work is devoted to chemical statics.

Production of a New Manure.—M. de Molon.—The author mixes finely-ground phosphate of lime with seaweeds, especially varec, and allows the mass to ferment for six to eight weeks.

Experiments on the Electric Discharge of the Chloride of Silver Battery.—MM. Warren de la Rue and H. W. Müller.—This paper has also appeared in English.

Action of Metallic Nitrates upon the Monohydrated Nitric Acid (see *Comptes Rendus*, lxxxix., 576).—A. Ditte.—There exists a group of nitrates which behave like magnesium nitrate; under the influence of heat they melt in their water of crystallisation which escapes along with the nitric acid, whilst a matter remains which still contains water in quantities more or less considerable, neutral nitrate, and either a sub-nitrate (zinc nitrate) or an oxide (manganese nitrate). In contact with the monohydrated acid the sub-salts are transformed into neutral nitrates, setting at liberty a certain quantity of water. The nitrates belonging to this group are those of magnesia, manganese, zinc, alumina, iron, copper, and uranium. Sub-nitrates are obtained from all of these metals except manganese, alumina, and iron. A third and numerous group, such as those of sodium, lithium, lime, barium, strontium, nickel, cobalt, bismuth, cadmium, mercury, silver, and lead, are insoluble or very sparingly soluble in the acid, at any temperature soever.

Silicium Nitride.—P. Schutzenberger.—The author finds that there exist two nitrides of silicon; the one Si_3N_2 , corresponding to cyanogen and titanium nitride, the other being very probably Si_3N_4 . The products resulting from the action of ammonia upon silicon chloride are again distinct, and contain either chlorine and hydrogen, or hydrogen alone. Their formulæ are $\text{Si}_3\text{N}_{10}\text{Cl}_3\text{H}$ and $\text{Si}_2\text{N}_3\text{H}$.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 2, September 11, 1879.

This issue contains no original chemical or physical matter. There is an article on the *Splendeurs de la Foi*,

and an attack upon the French Government for wishing to keep in its own hands the right of granting academical degrees.

No. 3, September 18, 1879.

Vines Reviving.—Certain vines at Neuvic (Charente Inférieure) supposed to be dead and about to be torn up have shot out afresh this year, and display a magnificent vegetation.

Trichinosis at Berlin.—Very stringent regulations have been adopted at Berlin to prevent the sale of trichinised pork. Every pig killed must be submitted to microscopic examination under a penalty of from 3 (sic) to 30 shillings.

No. 4, September 25, 1879.

This issue contains no matters of chemical interest.

No. 5, October 2, 1879.

Sophistication of Food and Drugs.—A statement is quoted from the *New York Popular Science Monthly* that a Boston chemist found in a sample of cream of tartar 75 per cent of *terra alba*, and that at Chicago, a chemist, in want of antimony sulphide, could find in the shops merely marble dust blackened with soot.

The Louis Maiche Battery.—The elements of this battery are platinised coke and amalgamated zinc, the latter placed in a solution of sal-ammoniac and the former plunging only slightly in the liquid, so as to present a large surface to the air.

Researches on the Movements of the Magnetic Needle.—E. Quetelet.—This memoir gives an account of the magnetic observations made at Brussels; a new study on the secular movements of the needle at the same locality, and a notice of the various movements which the needle experiences from different causes.

Researches on the Compressibility of Gases at High Pressures.—E. H. Amagat.—Already noticed.

Action of Ozone on the Colouring-matters of Plants.—A. Leeds.—Taken from the *CHEMICAL NEWS*.

New Method of Preparing Blesching Liquid.—Chloride of lime is decomposed with excess of bicarbonate of soda. The sediment subsides very easily.

No. 6, October 9, 1879.

Tempered Glass.—M. de la Bastie is said to have made great improvements both in the method of tempering glass and in the quality of the product. Among the articles shown to the Société d'Encouragement are mortars with their pestles for chemical use.

Production of Dew.—Prof. Levi Stockbridge.

Action of the Sun upon Galvanic Batteries.—Ph. Delahaye.—Already noticed.

Thermo-electric Battery.—M. Clamond.—A new battery for the production of the electric light. With a consumption of $9\frac{1}{2}$ kilos. coke, a current is produced capable of maintaining 4 lamps, each equal to 25 Carcel burners.

Researches with the Audiometer of Hughes.—B. W. Richardson.

Magnitude of Molecules.—D. C. Hodges.—These two papers are from an English source.

Influence of Atmospheric Electricity on the Growth, Inflorescence, and Fruification of Plants.—C. Naudin.—Already noticed.

Electro-motive Force in Free Jets of Water.—M. Elster.—Contrary to the assertions of Edlund, the friction of liquids against themselves produces no trace of electricity. If a jet of liquid is directed against a plate of some non-conducting substance, electric action is manifest. The currents observed in capillary tubes are due to the friction of the liquid against the sides.

Emission Spectra of the Haloid Compounds of Mercury.—B. Pierce.—On passing the spark into Geissler tubes containing a little of the salt and heating progressively, there is observed first the spectrum of the air, then that of the salt, and finally that of mercury. The chlorides give the same band between orange and green; mean wave-length $\lambda=443$, which seems to indicate a dissociation of HgCl_2 into HgCl and Cl . The bromide gives a band $\lambda=500$; the iodide, $\lambda=558$.

Influence of the Intensity of Sound upon the Speed of its Propagation.—H. Kayser.—The speed of sound in the free air is not affected by its intensity.

Electrolysis of Water.—M. Planer.—An electrolyte is always decomposed by a current, however feeble; but permanent decomposition only begins at the moment when the primary current is superior to the constant intensity of the polarisation current.

*Verhandlungen des Vereins zur Beforderung des
Gewerbflusses. No. 7, 1879.*

Chemistry at the Paris Exhibition.—Prof. C. Liebermann.—The author gives a review of the condensation of the so-called permanent gases; of the new element gallium, whereby he remarks that its identity with the theoretical "ekaaluminium" of Mendelejeff is not yet absolutely demonstrated; of the artificial jewels of Fremy and Feil, of Solvay's soda, the concentration of sulphuric acid, the fusion of platinum, tempered glass (the value of which he considers limited), the utilisation of kelp, lighting materials, the utilisation of beet-root treacle, and artificial colouring-matters. The industrial manufacture of artificial indigo he does not consider as near at hand.

*La Correspondance Scientifique.
No 65, October 21, 1879.*

Remedy against Rabies and the Bites of Serpents.—M. Torres Caicedo calls attention to the guaco and the cedron, two South American plants, as antidotes to the bites of serpents.

*Die Chemische Industrie.
No. 8, August, 1879.*

A considerable space is again devoted to a report of the debates in the German reichstag on the protective duties to be imposed upon chemicals.

Injury to Vegetation from Acid Vapours (Conclusion).—R. Hasenclever.—The author remarks that "in German towns which depend on the concourse of travellers, in watering places, &c., permission for the erection of industrial establishments is justly refused. [This is the principle which has been repeatedly advocated in the CHEMICAL NEWS that in certain districts the restrictions on the pollution, both of the atmosphere and of rivers, should be very severe, whilst in others considerable latitude may be allowed.] According to a private communication received by M. Hasenclever from Mr. Fletcher, the weekly quantity of noxious gases discharged into the atmosphere at St. Helens is as follows:—

Gaseous products of fuel	800 tons SO_2 .
From copper works	380 " "
From glass works	180 " "
From chemical works	25 " HCl .

In the neighbourhood of Ludwigshafen the plum trees are the only kind of vegetation apparently injured by the acid fumes.

*Reimann's Färber Zeitung,
No. 36, 1879.*

This issue contains nothing of general interest.

*Revue Universelle des Mines, de la Metallurgie, &c.,
Tome 5, No. 3, May and June, 1879.*

Study on the Influence of the Heat of the Interior of the Earth on the Possibility of Constructing Tunnels through High Mountains.—Dr. F. M. Stappf.—The author proposes the following questions:—(1.) At what degree of temperature will underground work be rendered impossible on physiological grounds? (2.) At what depth of a tunnel below the surface may the occurrence of such a temperature be expected? After consulting Prof. Dubois Reymond and instituting experiments at his recommendation in the St. Gotthardt tunnel, the conclusion is drawn that it is possible to work for a short time at 60° in a dry atmosphere, but that if the air is saturated with moisture 40° is the limit. The author considers that the products of the explosion of dynamite in mines may include nitric acid and carbonic oxide. Particles of nitroglycerine are also held in suspension in the smoke, and occasion head-ache, indigestion, and other distressing symptoms. The second question will be discussed in a continuation of this metal.

*Bulletin de la Société d'Encouragement pour l'Industrie
Nationale.
No. 68, August, 1879.*

This issue contains no chemical matter, and is chiefly devoted to an account of the calculating machine (arithmometer) of M. Thomas.

*Justus Liebig's Annalen der Chemie,
Band 198, Heft 3.*

Action of Elevated Temperatures and the Vapours of Carbolic Acid upon Organic Bodies.—Carl von Than.—During the outbreak of the plague in Russia the Hungarian sanitary authorities required that all substances imported from suspicious districts should be submitted to the joint action of high temperatures and the vapours of phenol, and the author was officially instructed to examine experimentally the action of this process of disinfection. He found that the conduction of heat through packets of letters, &c., was very slow. A packet of 22 letters was placed in an air-bath maintained at 111° . After four hours the temperature in the middle of the packet did not exceed 100° , whilst in a larger packet of 70 letters similarly treated the heat only rose to 75° . For practical purposes the letters were laid loosely in a wire basket, with a sheet of tin-plate or wire gauze between every ten, and in this manner any number of letters could be brought in two hours to the temperature of 148° . The second question was, What is the highest temperature which organic matter, especially paper, can bear without injury? It was found that in an air-bath, where the action of radiant heat was excluded, decomposition set in between 150° to 160° , but that at 140° no essential change took place. If exposed to the vapours of crystallised phenol at temperatures of from 130° to 140° , neither the paper nor the writing was injured. Filter and litmus paper, cotton, various kinds of linen, white and dyed tissues of wool and silk, lace, raw wool, and leather were not injured, either by the action of heat alone or by exposure to phenol vapours at 138° . Exceptions must be admitted in case of white wool, which was turned rather yellowish, and of wash-leather, which condensed a considerable quantity of phenol and became brittle. The action of heat alone and conjointly with phenol vapours was next studied. It was found that a dry heat of 137° delayed putrefaction, but did not suffice for the total destruction of all kinds of Bacteria. But a temperature of 137° applied along with the vapours of phenol completely eradicated all living organisms. These conclusions, therefore, confirm the experimental results of the late Prof. Crace-Calvert.

Determination of the Soluble Phosphoric Acid in Superphosphates.

I. Commercial Superphosphates.—E. Wein, L. Roesch, and J. Lehmann.—The process of determining phosphoric acid as agreed upon in 1872 by the agricultural chemists of North Germany, at the Magdeburg conference, having been called in question, the authors have submitted it to re-examination, and consider that the variation in its results is mainly due to differences in the method of preparing an aqueous extract of the sample. The process adopted at the Conference is as follows:—20 grms. superphosphate are stirred up with water in a mortar; the lumps broken without too great pressure, and the whole rinsed into a litre flask. So much water is then added that the liquid can be conveniently shaken, and digested for a time. If little iron and alumina is present the digestion was prolonged for two hours, but if these impurities were abundant the liquid was filtered off immediately after a careful agitation. After a very extensive series of experiments, the authors conclude that a digestion of two hours is not injurious in presence of ferric and aluminous phosphates; that the lixiviation of the sample upon a filter with the aid of the Bunsen pump offers no advantages over the method above described, and that the quantity of water generally used (1 litre to 20 grms.) is perfectly sufficient.

II. Superphosphates from Chemically Pure Material.—E. Wein.—The principal results are:—In superphosphates containing much free phosphoric acid, a very brief digestion is sufficient. In such cases the lixiviation process gives accurate results, though it is not to be recommended. Superphosphates containing but little free phosphoric acid should be digested for two hours to make sure that all soluble phosphates are extracted. In such cases the lixiviation method is not applicable. Superphosphates containing no free phosphoric acid must likewise be digested for two hours with a litre of water, but a further increase of the quantity does not seem serviceable.

Separation and Determination of Manganese.—J. Volhard.—(See p. 207.)

Heptan from Pinus Sabiniana.—T. E. Thorpe.

Derivatives of Isodurol.—Max Bielefeldt.—Not adapted for abstraction.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin. No. 11, 1879.

Separation of Ortho-xylol from its Isomers, and on a New Xylidin.—E. Wroblevsky.—Xylol is nitrised and reduced, yielding a mixture of the isomeric xylidins, the derivatives of meta- and ortho-xylol. The mixture is heated for three days with an equivalent quantity of glacial acetic acid. On submitting the mass to fractionated distillation the portion passing over up to 310° consists of water, acetic acid, and unchanged xylidin acetate. Above 320° acetxylid distils over, fusible at 127°. It is a derivative of metaxylol. The portion which went over below 310° is treated with soda-lye; when much unchanged xylidin floats upon the surface, which is distilled over with watery vapour and again heated with glacial acetic acid, after which it no longer congeals. This portion is again submitted to distillation when the derivative passing over above 320° crystallises readily, and consists of a mixture of acetxylids. The xylidin separated from the portion boiling at a lower temperature was found to be a derivative of orthoxylol.

On Protagon.—A. Gamgee and E. Blankenhorn.—The authors have been able to confirm the statements of Liebreich, both as regards the chemical and physical properties of protagon.

Formyl-tricarbonic Ester.—M. Conrad.—Formyl-tricarbonic ethyl-ester is a colourless liquid of pleasant odour, insoluble in water, volatile between 254° to 260°, of sp. gr. 1.10 at 19°, and having the composition $C_{10}H_{16}O_6$.

Reply to H. Kraut.—H. Kœbler.—The author maintains the accuracy of his observations on the melting-point of mercuric chloride.

Ellagic Acid.—L. Barth and G. Goldschmiedt.—In this very extensive memoir the authors endeavour to ascertain the constitution of ellagic acid and for this purpose they examine its behaviour with lime or soda-lime, with hydriodic acid, with hydrochloric acid, with caustic potassa, and with caustic soda. Summing up all their observations they think ellagic acid ought to be called "hexa-oxidiphenyl-keton-carbonic anhydride."

Action of Melting Soda upon Aromatic Acids.—L. Barth and J. Schröder.—As a rule these acids, when treated with melting potassa, are decomposed with elimination of carbonic acid. It was not found possible in this manner to arrive at more highly hydroxylised derivatives, as is the case with certain phenols.

Derivatives of α -Disulpho-phenolic Acid.—L. Barth and M. v. Schmidt.—Among the results obtained is the preparation of the sodium and the barium salts.

Disulpho-resorcinic Acid.—V. Tedeschi.—This acid is soluble in water and alcohol, insoluble in ether. The aqueous solution yields a splendid ruby-red colouration with ferric chloride. Its composition, according to the author, is $C_6H_2(SHO)_2OH_2 + 2H_2O$.

Action of Phosphorus Penta-chloride upon Saccharic Acid and Saccharine Bodies.—C. J. Bell.—After examining the behaviour of phosphorus penta-chloride upon saccharic acid, mannit, and dulcitol, the author concludes that the formation of chlor-muconic acid is no isolated fact, and that the isomeric saccharic acid and the corresponding kinds of sugar show a concordant behaviour.

A Correction.—S. Hoogewerff and W. A. van Dorp.—The authors used 8.5 to 9.5 grms. potassium permanganate not to 16 grms., but to 1 gram. dry quinine sulphate. See *Berichte*, xii., p. 158.

Constitution of Anthrarufin and Oxy-anthrarufin.—C. Liebermann and J. Dehnst.—Not suitable for abstraction.

Exsiccator for Carbon Disulphide, Ether, Chloroform, and Benzol.—C. Liebermann.—For the evaporation of these solvents the author charges a common exsiccator with fragments of crude paraffin instead of sulphuric acid.

Certain Derivatives of Ortho-nitranilin.—Ch. Rudolph.—The author dissolved ortho-nitranilin in chloroform, and heated the solution in a cohobator along with chloro-carbonic ether. Among the products formed was ortho-nitrophenyl-urethan.

On Benzylamin.—Ch. Rudolph.—A preliminary notice on the derivatives of benzylamin.

On Chlorinised Metallacetic Ethers.—F. Allihn.—An account of the preparation and properties of cuproacetmono-chloracetic ether.

A New Double Salt of Chromic Acid.—C. Hengen.—On adding a concentrated solution of neutral potassium chromate to one of ferric chloride as long as a precipitate is formed, and dissolving the latter in the necessary quantity of hydrochloric acid, a deep red solution is obtained, which in a few days deposits crystals of the new compound, $K_2CrO_4 \cdot Fe_2(CrO_4)_3 \cdot 4H_2O$.

On Para-diethyl Benzol from Para-dibrom Benzol.—H. Aschenbrandt.—An account of mono-sulpho-para-diethyl benzoic acid and mono-nitro-para-ethyl benzoic acid.

Action of Organic Zinc Compounds upon Quinones.—F. R. Japp.—In this paper the author describes the action of zinc-ethyl upon phenanthren-quinone.

On a Remarkable Decomposition of Phenyl-ethylamin Hydrochlorate.—M. Fileti and A. Piccini.—If the hydrochlorate of phenyl-ethylamin is heated to a

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THE CHEMICAL NEWS.

VOL. XL. No. 7041.

A NEW PROCESS IN METALLURGY.

By JOHN HOSKIN.

THE theory of this process has been developed from known principles, aided by experimental work undertaken for the purpose of investigating the action of rapid oxidation upon pyritous substances, with a view to their metallurgic treatment upon a large scale. Before these experiments were made metallurgists had not realised the fact that pyrites and other sulphides (even with the addition of a considerable proportion of incombustible materials) can be decomposed and fused by the heat developed in the oxidation which takes place whenever air is rapidly brought into contact with an excess of molten sulphides. When this is effected by introducing air under pressure through apertures of a few millimetres in diameter in the bottom of a hearth upon which the molten sulphides lie, the results produced are very remarkable. Thus, when cupreous pyrites was so treated, a true combustion of the more oxidisable constituents took place, flame and incandescence resulted, and the decomposition was effected with great rapidity.

It was primarily surmised that in this manner, neglecting the influence of mass, the elements would be burnt in the regular order of their relative affinities for oxygen, and that the second atom of sulphur in iron pyrites, which can be expelled by fusion, would escape oxidation in the molten bath and be volatilised in the current of sulphurous acid and nitrogen emerging from the surface of the molten liquid. The more volatile oxides and sulphides in the material operated upon, such as those of arsenic, antimony, lead, and zinc, would volatilise with this freed sulphur, and condense partly before it, though more or less contaminating that product. If the oxidation be arrested at a point determinable by practice, calculation, or some marked change in the spectrum, two products of different specific gravity will be obtained, namely, a slag of silicate of iron, lime, alumina, &c., containing the iron protoxide resulting from the oxidation of the iron sulphide, combined as silicate with the silicious fluxing materials present in the bath; and underneath this, the regulus or remaining unburnt sulphides, containing in an approximately known state of concentration the more valuable metals derived from the metalliferous substances operated upon. It was, however, necessary for the practical application of the theory that sufficient heat should be developed during the operation. The temperatures of combustion of various sulphides, calculated from known data, approached the maximum temperature attainable by the combustion of coal, and this inspired a considerable amount of confidence. In the case of iron pyrites these calculations are only rough approximations, as the latent heat of sulphur vapour is not known. It was found that when thus treating cupreous pyrites, the order in which the elements became oxidised was as follows:—

- | | | |
|----------------|----------|------------------|
| I. | II. | III. |
| Zinc and iron. | Sulphur. | Lead and copper. |

The above reactions find a parallel in the elimination of the metalloids from cast-iron by Bessemer's process, in which silicon and carbon and then phosphorus and manganese are successively burnt out of the crude metal. Parallel analogies also exist between the processes of puddling and English copper smelting; where the oxidation pro-

ceeds but slowly, and the necessary heat is obtained by the burning of coal.

The foregoing conclusions have been verified experimentally; full particulars thereof will be found in papers brought before the Society of Arts, February 12 and April 30 of this year. Without recording the crucible experiments, I will proceed to describe briefly those made at Penistone and Sheffield. In the former place, ordinary Bessemer converters were used, as they were the only existing plant available, but it was from the first contemplated that a very different description of furnace would be requisite. The first of these experiments was made on the night July 10 of last year, at Messrs. Charles Cammell's Penistone Works. The pyrites employed contained about 2½ per cent copper, 1½ ozs. silver, and approximately 3 grs. of gold per ton. About 6 tons of this mineral was melted in a cupola furnace, and the resulting proto-sulphides run into a converter. In this instance no silicious fluxes were added, the requisite silica being obtained from the gannister lining. The blast of air was at a pressure of from 15 to 20 lbs. The temperature gradually but steadily increased; a flame burnt continuously at the mouth of the vessel, where, after about twenty minutes' blow a pyrometer marked 1000° C., and wrought-iron melted in the incandescent sulphur vapour. The contents of the vessel were now very liquid, and at this period the protoxide of iron appears to have become converted into silicate as soon as it was formed. After thirty minutes' blow the silicious lining gave way at a weak place, cut through by the protoxide of iron, and a thin stream of very hot and liquid slag poured into the pit below. The converter was therefore emptied. The slag, consisting of ortho-silicate of iron, 2FeO.SiO₂, crystallised on cooling in large and well defined crystals above a regulus of iron and copper, the line of demarcation between the two being very distinct. If the oxidation had been continued longer a yet higher temperature would have been obtained. On the succeeding night the experiment was repeated several times, but silicious material was thrown into the vessel during the operations, which greatly mitigated the corrosive action of the protoxide of iron on the silicious lining. On 17-18th July further experiments were made, and valuable data obtained. The pyrites used was the same as before. Red sand was thrown into the converters.

In experiments made at the commencement of November, 1878, the blow, after having been commenced with some molten proto-sulphide, was continued with the addition of 4 tons of cold pyrites in large lumps, together with 9 cwt. of moist sand. When the converter became too full of material, half the charge was tipped out, and the blow was continued with the addition of 18 cwt. of pyrites and 3 cwt. of sand. On the 5th of February last about 18 tons of raw pyrites, with sandstone, were treated continuously in this manner in one Bessemer, being converted into silicate of iron slag and a rich regulus at the rate of over 2 tons an hour. In all these experiments the blast entered cold and the gases escaped at a high temperature; sufficient quantities of silicious fluxes were not added, and the resulting dense slag occasionally entrapped considerable quantities of copper. But in specially constructed furnaces, where the heat produced is properly taken advantage of, and possibly augmented, much larger quantities of flux can be added, which will enable a slag to be produced containing less copper than is now found in the rejected slags of the Swansea process.

Further experiments were made at Messrs. John Brown and Co.'s Atlas Works, Sheffield. A rough plant was extemporised by connecting several ordinary cupolas, as shown in the diagram, one of which was provided with a Bessemer hearth. The sulphur and sublimates were condensed in the second and third cupolas. A large portion of the theoretical yield of sulphur was thus obtained, notwithstanding the imperfect appliances used. Slag and regulus were produced as before, and it was evident that the process could be carried on continuously. These ex-

* Read before the British Association for the Advancement of Science (Section B.), Sheffield, 1879.

periments were not continued for any length of time, partly owing to the faulty construction of the forepart of the furnace, and partly to the irregular manner in which the materials were introduced.

The spectroscopic observations taken by Dr. W. M. Watts during the course of these experiments were valuable and interesting, and I am indebted to him for the following information:—

In the experiments at Penistone two spectra were observed; the first, that given by the flame from the charging-door of the cupola in which the pyrites was melted; the second, produced by the blast of air through the molten protosulphide in the converter. The cupola-spectrum was shown by direct comparison with the spectrum of a flame coloured by lead chlorate to be mainly due to oxide of lead, but contained besides some few of the lines which appear to be proper to the converter-spectrum. Analysis showed that the lead present in the ore was almost entirely volatilised during the preliminary melting of the ore, the molten protosulphide charged into the converter containing only 0.8 per cent lead. The converter flame gives a brilliant spectrum extending from the lithium line somewhat beyond the thallium line, which is usually present. Its most marked feature is the presence of four bright red lines about equally spaced between the sodium and lithium lines. Their wave-lengths, as far as at present known, are approximately 5999, 6151, 6320, and 6466, besides a fainter line at 6113. These lines are not, those of any known spectrum. The way in which the flame is obtained suggests the theory that they are sulphur lines. When sulphur is burned in air or oxygen the spectrum obtained is entirely continuous, and even if air be bubbled through boiling sulphur no lines are obtained. Two spectra of sulphur obtained by the electric discharge through a vacuum tube containing vapour of sulphur have been described, but neither contains these four red lines. The spark with a Leyden jar in a current of sulphur dioxide at the ordinary pressure yields a spectrum (at present under investigation) apparently not previously described, in which, however, the red lines are altogether different from those of the converter-spectrum. The constancy with which these four red lines are associated together seems to preclude the possibility of their being due to different substances, otherwise the most refrangible line might be due to lead. No lines of copper were observed except in the fourth experiment, in which all the lines except those of sodium disappeared about six minutes before the turn down. When in this experiment, towards the end of the blow, the sub-sulphide of copper began to burn, a splendid emerald green flame suddenly appeared, and all the lines except those of copper and sodium left the spectrum. During the last few minutes of the blow the mouth of the converter was dull and without flame, the sulphur and oxidisable matter having been burnt out.

The pyrites, after fusion in the cupola at Penistone, gave protosulphides containing:—

	I.* Per cent.	II.* Per cent.
Iron	59.62	60.30
Copper	3.52	3.25
Zinc	1.52	1.88
Lead	0.79	0.81
Arsenic	0.06	0.05
Manganese	0.21	0.20
Alumina	0.15	trace
Lime	0.28	0.34
Magnesia	0.27	0.32
Sulphur	33.10	32.50
Silica	0.15	0.30
	99.67	99.95

The products consisted in the first place of a regulus having a greater density than ordinary copper regulus on account of the larger proportion of iron, as compared with sulphur therein contained. The average specific gravity is 4.96. The following analyses illustrate the composition:—

	I.* Per cent.	II.* Per cent.	III.† Per cent.	V.‡ Per cent.	VI.‡ Per cent.
Iron	57.10	56.05	12.56	13.20	13.16
Copper	15.85	16.59	62.36	63.43	59.98
Zinc	0.84	0.48	0.42	Silver and	0.152
Lead	0.22	0.31	0.14	Gold	
Arsenic	0.04	0.03	—	—	—
Manganese	0.22	0.20	—	—	—
Alumina	0.11	0.13	—	—	—
Lime	0.34	0.16	—	—	—
Magnesia	0.34	0.25	—	—	—
Sulphur	21.96	23.47	22.22	20.37	21.94
Silica or insoluble residue ..	2.00	1.10	0.28	1.20	2.57
Oxygen and not estimated ..	0.98	1.23	2.02	1.80	2.198
	100.00	100.00	100.00	100.00	100.000

	I.* oz. dwt. gr.	II.* oz. dwt. gr.	V.‡ oz. dwt. gr.	VI.‡ oz. dwt. gr.
Silver per ton of regulus	10 16 10	11 15 10	54 1 6	48 5 6
Gold ditto ..	0 6 12	0 7 0	1 1 5	1 4 11

The slag, the second product, when produced without the use of extraneous basic flux, is a black, lustrous, normal silicate of protoxide of iron, having the formula $2\text{FeO}, \text{SiO}_2$. Sulphur sometimes replaces part of the oxygen in it. Some specimens were very finely crystallised; the specific gravity is 4.05. This slag keeps very liquid at the temperature of the operation. The following represents the normal composition when using only sand as flux, and when no precautions are taken to obtain a complete separation of the regulus from the slag:—

	I.* Per cent.	II.* Per cent.	III.† Per cent.	V.‡ Per cent.	VI.‡ Per cent.
Iron protoxide ..	53.30	54.62	59.00	67.17	67.52
Iron peroxide ..	3.00	3.71	—	—	—
Iron combined with sulphur	5.79	4.27	—	—	—
Copper ditto ..	0.16	0.22	1.55	0.87	0.42
Lead	0.12	0.10	—	—	—
Zinc oxide	1.15	1.75	0.98	—	—
Arsenic	trace	trace	none	—	—
Manganese oxide	0.32	0.37	0.30	—	—
Alumina	2.15	2.06	1.08	—	2.46
Lime	0.40	0.37	0.63	0.07	—
Magnesia	0.46	0.45	—	—	—
Sulphur	3.39	2.55	6.67	1.08	2.06
Silica	29.90	30.05	29.55	28.53	26.22
Oxygen and not estimated ..	—	—	0.24	2.28	1.32
	100.14	100.52	100.00	100.00	100.00

When, however, the products are overblown the proto-silicate burns to silicate of peroxide of iron, a more infusible and less dense silicate, of which the following is an analysis:—

	IV.† Per cent.
Silica	34.34
Iron protoxide	25.10
Iron peroxide	33.83
Manganese protoxide	0.12
Alumina	1.81
Zinc oxide	0.73
Copper oxide	2.39
Lead oxide	0.03
Lime	0.24
Magnesia	0.30
Sulphur	0.15
Arsenic	none
Phosphoric acid	0.031
Not estimated and insoluble residue	1.45
	100.521

* Analysed by Mr. J. E. Stead. † By Mr. E. Riley.
‡ By Mr. A. E. Arnold.

The third class of these products consists of metallic sublimates, and of sulphur. As an example of the former the following may be quoted:—

	Per cent.	
Sulphate of lead	52.08	} 50.91 % Pb.
Sulphide	17.29	
Zinc oxide	21.78	} 17.47 „ Zn.
Copper oxide	0.09	
Iron sesquioxide	2.86	} 0.07 „ Cu.
Insoluble residue	2.14	
Not estimated and loss..	3.76	
	100.00	

The following is an analysis by Mr. J. S. Merry, of Swansea, of the crude sulphur dried at 100° C.:—

	Per cent.
Sulphur	63.30
Silica	7.00
Iron	2.76
Arsenic	7.12
Zinc	2.74
Lead	8.95
Copper	trace
Lime	trace
Magnesia	trace
Alumina.. ..	trace
Oxygen (traces), carbon and loss ..	8.13
	100.00

The fourth, the gaseous product of the operation, consists mainly of sulphurous acid and nitrogen. The following are the analyses by Dr. Frankland, F.R.S.:—

	I. Per cent.	II. Per cent.
Nitrogen	86.00	88.37
Sulphurous acid	14.00	10.88
Oxygen	—	0.75
	100.00	100.00

The principal cost of plant will be for the blast. I am indebted to Mr. W. H. Cutler, C.E., Queen Anne's Gate, Westminster, for the following information:—"The cost of a suitable pair of air pumps to pump 2000 tons of air per month of 28 days, working day and night, or 24,000 tons of air per annum, each pump capable on emergency of pumping the required quantity of air at 10 lbs. pressure on the square inch, viz., 30½ inches diameter with 5 feet stroke, on strong cast iron bed-plates, will cost £742 10s. The power required will be 100 horse-power. The dimensions and cost of the turbine must depend on the fall of water. For example, a fall of 72 cubic feet per second for 18 feet would drive a turbine 79 revolutions per minute, and produce 108 horse-power." The cost of one of Mr. Cutler's turbines, of suitable dimensions, would be £220. That of the furnace would not exceed £100, and say £200 for a culvert for collecting the sulphur and other sublimates. Twenty-four thousand tons of air would be more than sufficient to supply the necessary oxygen to 15,000 ton of pyrites, since 1.4 part by weight of air will convert one part of ordinary cupreous pyrites into rich regulus. It therefore appears from the foregoing figures that where sufficient water power is available a plant capable of treating 15,000 tons of pyrites annually could be erected at a cost of about £1500. Where, however, water power is not available steam boilers will be requisite, and the additional cost for plant may be £500 or perhaps £1000. Messrs. Howson and Wilson estimate that com-

pound engines of 2400 horse-power, with 23 boilers of ordinary size heated by 20,000 tons of coal, would be sufficient to supply 480,000 tons of air annually at a pressure of 17 pounds per square inch. This quantity is more than sufficient to supply the necessary oxygen to 300,000 tons of pyrites. The same gentlemen estimate that the consumption of coal requisite to heat the blast to 1000° F. would be about 11,000 tons per annum, but this quantity will be materially decreased if the heat from the exit gases be utilised.

With regard to the furnace it is proposed to make the hearth, or rather crucible, of silicious, aluminous, or refractory carbonaceous material. A sufficiently large proportion of silicious flux in the furnace charge will greatly mitigate the action of the resulting iron protoxide upon the silica of the lining. Aluminous shrunk bricks may answer still better. It might even be found convenient to allow considerable corrosion to the lining to take place, if the converting hearth is of such form, and the materials are of such a nature, that it can be readily and economically renewed.

It may be also advantageous to run the regulus and slag, after the desired concentration has been effected, directly on to the hearth of a reverberatory furnace, where they can be kept molten by external heat, and where a more perfect separation of the one from the other may be effected. In such a furnace the final oxidation of the rich regulus would probably be most conveniently effected, although it is of course possible to produce metallic copper from the regulus by the transmission of air currents in a specially constructed furnace.

Not only would antimony, lead, zinc, copper, nickel, silver, and other valuable metals be extracted from the sulphides that contain them, but also from the incom-bustible fluxing materials that are added to the charge, and the extraction of the copper, and silver and gold will probably be more complete, than by any known process. In countries where cupreous silicious schists and sand-stones abound, the use of these as silicious fluxes would partially, if not wholly, compensate for the loss of copper in the slag. Thus, by using 0.5 ton of such material, containing 0.5 per cent of copper for each ton of the sulphuretted ore, the whole of the copper could be recovered from the latter, assuming the slag to contain even as much as 0.2 per cent of that metal.

The crude sulphur may be freed from the accompanying sulphide of arsenic by boiling it with milk of lime, and from the metallic oxides and sulphides with which it is contaminated by distillation; or purification by bisulphide of carbon might be resorted to. The sulphurous acid can be oxidised in chambers to sulphuric acid, either with or without previous liquefaction.

This process, on account of its simplicity and economy, may reasonably be expected, not only to take the place of the ordinary smelting, but also of many of the wet processes now in use.

Examination and Estimation of Manurial Matters especially Bone Dust.—Prof. Krockner.—Ground bones have often been previously completely exhausted of gelatin and horn dust, leather waste, &c., are afterwards added to make up the guaranteed proportion of nitrogen. Good ground bones should contain to 20 to 22 per cent of phosphoric acid at least 3 to 3½ per cent of nitrogen due to gelatin, to which even in the best samples horn dust is added so as to bring up the total nitrogen to 3½ to 4 per cent. To ascertain the presence of such additions the author shakes up bone meal with chloroform, when horn, blood, leather waste, &c., rise to the surface, whilst bone earth mixed with its natural gelatinous matter, and pure phosphates remain at the bottom. So-called vegetable ivory remains also at the bottom, and must be recognised microscopically. Horn dust as met with in commerce contains about 15 per cent of nitrogen.—*Biedermann's Central-blatt für Agrikultur. Chemis.*

* Analysed by Mr. A. E. Arnold.

ON THE VAPOUR-DENSITIES OF
PEROXIDE OF NITROGEN, FORMIC ACID,
ACETIC ACID, AND PERCHLORIDE OF
PHOSPHORUS.

By J. WILLARD GIBBS.

THE relation between temperature, pressure, and volume, for the vapour of each of these substances differs widely from that expressed by the usual laws for the gaseous state—the laws known most widely by the names of Mariotte, Gay-Lussac, and Avogadro. The density of each vapour, in the sense in which the term is usually employed in chemical treatises, *i.e.*, its density taken relatively to air of the same temperature and pressure,* has not a constant value, but varies nearly in the ratio of one to two. And these variations are exhibited at pressures not exceeding that of the atmosphere and at temperatures comprised between zero and 200° or 300° of the centigrade scale.

Such anomalies have been explained by the supposition that the vapour consists of a mixture of two or three different kinds of gas or vapour, which have different densities. Thus, it is supposed that the vapour of peroxide of nitrogen is a gas-mixture, the components of which are represented (in the newer chemical notation) by NO_2 and N_2O_4 respectively. The densities corresponding to these formulæ are 1.589 and 3.178. The density of the mixture should have a value intermediate between these numbers, which is substantially the case with the actual vapour. The case is similar with respect to the vapour of formic acid, which we may regard as a mixture of CH_2O_2 (density 1.589) and $\text{C}_2\text{H}_4\text{O}_4$ (density 3.178), and the vapour of acetic acid, which we may regard as a mixture of $\text{C}_2\text{H}_4\text{O}_2$ (density 2.073) and $\text{C}_4\text{H}_8\text{O}_4$ (density 4.146). In the case of perchloride of phosphorus, we must suppose the vapour to consist of three parts:— PCl_5 (the proper perchloride, density 7.20), PCl_3 (the protochloride, density 4.98), and Cl_2 (chlorine, density 2.22). Since the chlorine and protochloride arise from the decomposition of the perchloride, there must be as many molecules of the type Cl_2 as of the type PCl_3 . Now a gas mixture containing an equal number of molecules of PCl_3 and Cl_2 will have the density $\frac{1}{2}(4.98 + 2.22)$ or 3.60. It follows that, at least so far as the range of the possible values of its density is concerned, we may regard the vapour as a mixture in variable proportions of two kinds of gas having the densities 7.20 and 3.60 respectively. The observed values of the densities accord with this supposition.

These hypotheses respecting the constitution of the vapours are corroborated, in the case of peroxide of nitrogen and perchloride of phosphorus, by other circumstances. The varying colour of the first vapour may be accounted for by supposing that the molecules of the type N_2O_4 are colourless, while each molecule of the type NO_2 has a constant colour. This supposition affords a simple relation between the density of the vapour and the depth of its colour, which has been verified by experiment.†

The vapour of the perchloride of phosphorus shows with increasing temperature in an increasing degree the characteristic colour of chlorine. The amount of the colour appears to be such as is required by the hypothesis respecting the constitution of the vapour on the very probable supposition that the perchloride proper is colourless, but the case hardly admits of such exact numerical determinations as are possible with respect to the peroxide of nitrogen.‡ But since the products of dissociation are in this case dissimilar, they may be partially separated by diffusion through a neutral gas, the lighter chlorine diffusing more rapidly than the heavier protochloride. The

fact of dissociation has in this way been proved by direct experiment.*

In the case of acetic and formic acids, we have no other evidence than the variations of the densities in support of the hypothesis of the compound nature of the vapour, yet if these variations shall appear to follow the same law as those of the peroxide of nitrogen and the perchloride of phosphorus, it will be difficult to refer them to a different cause.

But however it may be with these acids, the peroxide of nitrogen and the perchloride of phosphorus evidently furnish us with the means of studying the laws of chemical equilibrium in gas mixtures in which chemical change is possible and does in fact take place, reversibly, with varying conditions of temperature and pressure. Or, if from any considerations we can deduce a general law determining the proportions of the component gases necessary for the equilibrium of such a mixture under any given conditions, these substances afford an appropriate test for such a law.

In a former paper† by the present writer equations were proposed to express the relation between the temperature, the pressure, or the volume, and the quantities of the components in such a gas mixture as we are considering—a gas mixture of convertible components in the language of that paper. Applied to the vapour of the peroxide of nitrogen, these equations led to a formula giving the density in terms of the temperature and pressure, which was shown to agree very closely with the experiments of Deville and Troost, and much less closely, but apparently within the limits of possible error, with the experiments of Playfair and Wanklyn. Since the publication of that paper, new determinations of the density have been published in different quarters, which render it possible to compare the equation with the results of experiment throughout a wider range of temperature and pressure. In the present paper all experimental determinations of the density of this vapour which have come to the knowledge of the writer are cited, and compared with the values demanded by the formula, and a similar comparison of theory and experiment is made with respect to each of the other substances which have been mentioned.

The considerations from which these formulæ were deduced may be briefly stated as follows. It will be observed that they are based rather upon an extension of generally acknowledged principles to a new class of cases than upon the introduction of any new principle.

The energy of a gas mixture may be represented by an expression of the form—

$$m_1(c_1t + E_1) + m_2(c_2t + E_2) + \&c.,$$

with as many terms as there are different kinds of gas in the mixture, $m_1, m_2, \&c.$, denoting the quantities (by weight) of the several component gases, $c_1, c_2, \&c.$, their several specific heats at constant volume, $E_1, E_2, \&c.$, other constants, and t the absolute temperature. In like manner the entropy of the gas mixture is expressed by—

$$m_1 \left(H_1 + c_1 \log x t - a_1 \log x \frac{m_1}{v} \right) \\ + m_2 \left(H_2 + c_2 \log x t - a_2 \log x \frac{m_2}{v} \right) + \&c.$$

Where x denotes the volume, and $H_1, a_1, H_2, a_2, \&c.$, denote constants relating to the component gases, $a_1, a_2, \&c.$, being inversely proportional to their several densities. The logarithms are Napierian. These expressions for energy and entropy will undoubtedly apply to mixtures of

* Wanklyn and Robinson, "On Diffusion of Vapours: a means of distinguishing between apparent and real Vapour-densities of Chemical Compounds." *Proc. Roy. Soc.*, vol. xii., p. 507.

† "On the Equilibrium of Heterogeneous Substances." *Transactions of the Connecticut Academy*, vol. iii., p. 108. The equations referred to are (313), (317), (319), and (320) on pages 233 and 234. The applicability of these equations to such cases as we are now considering is discussed under the heading "Gas-mixtures with Convertible Components," page 234.

* The language of this paper will be conformed to this usage.

† Salet, "Sur la Coloration du Peroxyde d'azote." *Comptes Rendus*, t. lxviii., p. 488.

‡ H. Sainte-Claire Deville, "Sur les Densités de Vapeur." *Comptes Rendus*, t. lxvii., p. 1157.

different gases, whatever their chemical relations may be (with such limitations and with such a degree of approximation as belong to other laws of the gaseous state), when no chemical action can take place under the conditions considered. If we assume that they will apply to such cases as we are now considering, although chemical action is possible, and suppose the equilibrium of the mixture with respect to chemical change to be determined by the condition that its entropy has the greatest value consistent with its energy and its volume, we may easily obtain an equation between m_1 , m_2 , &c., t and v .*

The condition that the energy does not vary gives—

$$(m_1 c_1 + m_2 c_2 + \&c.) dt + (c_1 t + E_1) dm_1 + (c_2 t + E_2) dm_2 + \&c. = 0. \quad (1)$$

The condition that the entropy is a maximum implies that its variation vanishes when the energy and volume are constant: this gives—

$$\frac{m_1 c_1 + m_2 c_2 + \&c.}{t} dt + \left(H_1 - a_1 + c_1 \log n t - a_1 \log \frac{m_1}{v} \right) dm_1 + \left(H_2 - a_2 + c_2 \log n t - a_2 \log \frac{m_2}{v} \right) dm_2 + \&c. = 0. \quad (2)$$

Eliminating dt , we have—

$$\left(H_1 - a_1 - c_1 - \frac{E_1}{t} + c_1 \log n t - a_1 \log \frac{m_1}{v} \right) dm_1 + \left(H_2 - a_2 - c_2 - \frac{E_2}{t} + c_2 \log n t - a_2 \log \frac{m_2}{v} \right) dm_2 + \&c. = 0. \quad (3)$$

If the case is like that of the peroxide of nitrogen this equation will have two terms, of which the second may refer to the denser component of the gas mixture. We shall then have $a_1 = 2a_2$, and $dm_1 = -dm_2$, and the equation will reduce to the form—

$$\log \frac{m_2 v}{m_1^2} = -A - B \log t + \frac{C}{t}, \quad (4)$$

where common logarithms have been substituted for Napierian, and A , B , and C are constants. If, in place of the quantities of the components, we introduce the partial pressures, p_1 , p_2 , due to these components, and measured in millimetres of mercury, by means of the relations—

$$m_1 = \frac{p_1 v}{a_1 t}, \quad m_2 = \frac{p_2 v}{a_2 t},$$

where a_1 denotes a constant, we have—

$$\log \frac{p_2^2}{p_1^2} = -(A + \log 2a_1) - (1 + B) \log t + \frac{C}{t} \\ = -A' - B' \log t + \frac{C}{t}, \quad (5)$$

where A' and B' are new constants. Now if we denote by p the total pressure of the gas mixture (in millimetres of mercury) by D , its density relative to air of the same temperature and pressure, and by D_1 , the theoretical density of the rarer component, we shall have—

$$p : p + p_2 :: D_1 : D.$$

This appears from the consideration that $p + p_2$ represents what the pressure would become if, without change of temperature or volume, all the matter in the gas mixture could take the form of the rarer component. Hence—

$$p_2 = p \frac{D - D_1}{D_1},$$

$$p_1 = p - p_2 = p \frac{2D_1 - D}{D_1},$$

and—

$$\frac{p_2}{p_1^2} = \frac{D_1(D - D_1)}{p^2(2D_1 - D)^2}.$$

* For certain *a priori* considerations which give a degree of probability to these assumptions, the reader is referred to the paper already cited.

By substitution in (5) we obtain—

$$\log \frac{D_1(D - D_1)}{(2D_1 - D)^2} = -A' - B' \log t + \frac{C}{t} + \log p_1. \quad (6)$$

By this formula, when the values of the constants are determined, we may calculate the density of the gas mixture from its temperature and pressure. The value of D_1 may be obtained from the molecular formula of the rarer component. If we compare equations (3), (4), and (5) we see that—

$$B' = B + 1, \quad B = \frac{c_1 - c_2}{a_2}.$$

Now $c_1 - c_2$ is the difference of the specific heats at constant volume of NO_2 and N_2O_4 . The general rule that the specific heat of a gas at constant volume and per unit of weight is independent of its condensation would make $c_1 = c_2$, $B = 0$, and $B' = 1$. It may easily be shown, with respect to any of the substances considered in this paper,* that unless the numerical value of B' greatly exceeds unity, the term $B' \log t$ may be neglected without serious error, if its omission is compensated in the values given to A' and C . We may therefore cancel this term, and then determine the remaining constants by comparison of the formula with the results of experiment.

In the case of a mixture of Cl_2 , PCl_3 , and PCl_5 , equation (3) will have three terms distinguished by different suffixes. To fix our ideas, we may make these suffixes 2, 3, and 5, referring to Cl_2 , PCl_3 , and PCl_5 respectively. Since the constants a_2 , a_3 , and a_5 are inversely proportional to the densities of these gases—

$$a_2 dm_2 = a_3 dm_3 = -a_5 dm_5,$$

and we may substitute—

$$-\frac{1}{a_2}, \quad -\frac{1}{a_3}, \quad -\frac{1}{a_5}$$

for dm_2 , dm_3 , and dm_5 in equation (3), which is thus reduced to the form—

$$\log \frac{m_5 v}{m_2 m_3} = -A - B \log t + \frac{C}{t}. \quad (7)$$

If we eliminate m_2 , m_3 , m_5 by means of the partial pressures, p_2 , p_3 , p_5 , we obtain—

$$\log \frac{p_5}{4p_2 p_3} = -A' - B' \log t + \frac{C}{t} \quad (8)$$

when A' , B' , like A , B , and C , are constants. If the chlorine and the protochloride are in such proportions as arise from the decomposition of the perchloride, $p_2 = p_3$ and $4p_2 p_3 = (p_2 + p_3)^2$. In this case, therefore, we have—

$$\log \frac{p_5}{(p_2 + p_3)^2} = -A' - B' \log t + \frac{C}{t}. \quad (9)$$

It will be seen that this equation is of the same form as equation (5), when p_5 in (9) is regarded as corresponding to p_2 in (5), and $p_2 + p_3$ in (9), which represent the pressure due to the products of decomposition, is regarded as corresponding to p_1 in (5), which has the same significance. It follows that equation (5), as well as (6), which is derived from it, may be regarded as applying to the vapour of perchloride of phosphorus, when the values of the constants are properly determined. This result might have been anticipated, but the longer course which we have taken has given us the more general equations, (7) and (8), which will apply to cases in which there is an excess of chlorine or of the protochloride.

If the gas mixture considered, in addition to the components capable of chemical action, contains a neutral gas, the expression for the energy and entropy of the gas mixture should properly each contain a term relating to this neutral gas. This would make it necessary to add $c_n m_n$ to the coefficient of dt in (1), and $\frac{c_n m_n}{t}$ to the coefficient of

* For the case of peroxide of nitrogen see pp. 243, 244 in the paper cited above.

TABLE I.—PEROXIDE OF NITROGEN.
Experiments at Atmospheric Pressure.
MITSCHERLICH—R. MÜLLER—DEVILLE AND TROOST.

Tempera- ture.	Pres- sure.	Density calc. by eq. (10).	Density Observed. Deville and Troost.			Excess of Observed Density. Deville and Troost.			
			M—h. M—r.	I.	II.	III.	M—h. M—r.	I.	II.
183.2	(760)	1.592				1.57			—0.022
154.0	(760)	1.597				1.58			—0.017
151.8	(760)	1.598		1.50			—0.10		
135.0	(760)	1.607				1.60			—0.007
121.8	(760)	1.622		1.64			+0.02		
121.5	(760)	1.622				1.62			—0.002
111.3	(760)	1.641				1.65			+0.009
100.25	760	1.677	1.72				+0.04		
100.1	(760)	1.676				1.68			+0.004
100.0	(760)	1.677		1.71			+0.03		
90.0	(760)	1.728				1.72			—0.008
84.4	(760)	1.768		1.83			+0.06		
80.6	(760)	1.801				1.80			—0.001
79	748	1.814	1.84				+0.03		
77.4	(760)	1.833			1.85			+0.02	
70.0	(760)	1.920				1.92			0.000
70	754.5	1.919	1.95				+0.03		
68.8	(760)	1.937		1.99				+0.05	
66.0	(760)	1.976			2.03			+0.05	
60.2	(760)	2.067				2.08			+0.013
55.0	(760)	2.157			2.20			+0.04	
52	757	2.211	2.26				+0.05		
49.7	(760)	2.255		2.34				+0.09	
49.6	(760)	2.256				2.27			+0.014
45.1	(760)	2.342			2.40			+0.06	
39.8	(760)	2.443				2.46			+0.017
35.4	(760)	2.524				2.53			+0.006
35.2	(760)	2.528		2.66			+0.13		
34.6	(760)	2.539			2.62			+0.08	
32	748	2.582	2.65				+0.07		
28.7	(760)	2.642		2.80				+0.16	
28	751	2.652	2.70				+0.05		
27.6	(760)	2.661			2.70			+0.04	
26.7	(760)	2.676				2.65			—0.026

dt in (2), the suffix n being used to mark the quantities relating to the neutral gas. But these quantities would disappear with the elimination of dt , and equation (3) and all the subsequent equations would require no modification if only p and D are estimated (in accordance with usage) with exclusion of the pressure and weight due to the neutral gas. This result, which may be extended to any number of neutral gases, is simply an expression of Dalton's law.

We now proceed to the comparison of the formulæ, especially of equation (6), with the results of experiment.

(To be continued.)

ON ERBIA.

By P. T. CLÈVE.

AT the meeting of the Academy on September 15, M. Soret made some remarks on a memoir which I presented to the Academy on the 1st September, in which I stated that from an examination of the absorption spectra old erbium is capable of being divided into three different bodies. When I made the researches described in that memoir I was not aware of the work of M. Soret. From his remarks, however, it is evident that he had previously discovered the same facts. He consequently has the priority, and I am glad to have the opportunity of doing him justice.

The body for which I proposed the name of *holmium* is evidently the same that M. Soret calls X. I have not been able to identify it with the philippium of M. Delafontaine, because that body is characterised by an absorption ray

in the blue part of the spectrum, which ray occupies the same place as a ray of erbium. Besides, the body X possesses rather a high atomic weight, whilst that of philippium is pointed out as being very low. Anyhow, it has been proved, as a certain fact, that the rays $\lambda=640$ and $\lambda=536$ do not belong to erbium, as I have succeeded in almost entirely eliminating them from the spectrum of erbium. As to the body which I have called *holmium*, it is evident that M. Soret was again before me in observing the variations in intensity of its absorption ray. I think I have unmistakably proved that this ray does not belong to erbium, because I obtained a fraction of erbium whose spectrum did not contain that ray, and gave only traces of the rays of the body X. If this ray belongs neither to ytterbium nor to erbium, it seems to me that it must be due to some elements at present unknown. I am now engaged in procuring new materials for the preparation of these rare bodies, which are so difficult to separate. Very recently I began an experiment with 11 kilos. of gadolinite, the treatment of which is already so far advanced that it is to be hoped that the questions as to the yttria earths will soon receive their solution.—*Comptes Rendus*, October 27.

The Royal Polytechnic.—The Directors of this Institution are increasing the number of special daily entertainments. The more scientific portion of the new programme includes a description and exhibition of Edison's loud-speaking telephone which in itself forms a great attraction. There are also lectures on Chemistry of Coal and on Flashing Signals. A practical demonstration of Fleuss's system of Walking Under Water is also given by the inventor.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

OCTOBER, 1879.

THE following are the returns of the Society of Medical Officers of Health:—

	Appearance in a foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia	Chlorine	Sulphuric An- hydride.	Hardness on Clark's Scale		
		Saline.	Organic.								Before Boiling.	After Boiling.	
													Grains.
Thames Water Companies.													
Grand Junction	Clear	0'000	0'008	0'135	0'103	21'40	7'340	0'360	1'080	1'070	13'7	4'6	
West Middlesex	Clear	0'000	0'010	0'129	0'111	21'40	7'700	0'396	1'150	1'360	13'7	4'2	
Southwark and Vauxhall	Clear	0'000	0'008	0'117	0'088	21'60	7'280	0'468	1'150	1'300	13'2	4'2	
Chelsea	Clear	0'000	0'008	0'138	0'085	21'00	7'750	0'371	1'220	1'300	13'7	4'6	
Lambeth	Clear	0'000	0'009	0'148	0'092	21'10	7'530	0'428	1'220	1'300	14'8	3'3	
Other Companies.													
Kent	Clear	0'000	0'005	0'480	0'022	29'46	10'700	0'396	1'870	2'600	19'4	6'5	
New River	Clear	0'000	0'006	0'126	0'052	21'00	7'630	0'468	1'150	1'200	13'7	3'3	
East London	Clear	0'000	0'008	0'144	0'086	20'00	7'950	0'432	1'220	1'200	14'3	4'2	

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrites, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

ON THE
ESTIMATION OF CARBON IN CAST-STEELS.

By SERGIUS KERN, M.E., St. Petersburg.

THIS summer the author analysed several samples of crucible steel for carbon. In most of the laboratories of steel works Eggertz's method is used for that purpose. But as it was desired to estimate the contents of the carbon in the steels in hand as accurate as possible, the ordinary combustion method was adopted. Chromic acid was used. Meanwhile, all the samples were also tested by the Eggertz's process. The results of the analyses differ very much if steel is analysed by both methods. The following table gives the percentage of carbon found in the steels:—

Eggertz's Method.	Combustion Method.
0'12	0'14
0'15	0'17
0'24	0'27
0'34	0'38
0'36	0'37
0'45	0'46
0'60	0'64
1'03	1'02

In laboratories Eggertz's process may only be used for rough estimations, but it must be regretted that, at the present time, this rule is nearly abandoned.

CORRESPONDENCE.

EMPLECTITE.

To the Editor of the Chemical News.

SIR,—Having lately discovered at the Aamdal Copper Mines, Upper Thelemarken, Norway, the mineral emplectite, and observing that Dana in his system of mineralogy, does not mention any localities in Norway for this mineral, I gladly hand you this notice of its discovery here for your valuable journal.

The mineral, on analysis, gave,—

Bismuth	57'72
Copper	17'23
Silver	2'91
Lead	a trace
Sulphur	19'20
Silica	1'30
	98'36

The silver here without doubt replacing a part of the bismuth or copper, allowing the composition of this mineral to be of the formula $\text{CuS} + \text{Bi}_2\text{S}_3$ —I am, &c.,—

F. R. W. DAW.

Aamdals Kobberværk, Skafse,
Ovre Thelemarken.
October 25, 1879.

THE DISSOCIATION OF CHLORINE.

To the Editor of the Chemical News.

SIR,—If Mr. F. P. Dunnington had not the impression that PtCl_4 (platonic chloride) was the source of the chlorine with which Victor Meyer performed his experiments, why base suppositions or suggestions of error upon reactions of a compound which was not used? The residue Mr. Dunnington obtained on heating this, consisting, as he says, "of PtCl_2 , with but little Pt and PtCl_4 ," is the real substance in question; and as nothing is said of oxygen compounds in this, Mr. Dunnington had surely nothing on which to base the question of "due care having been taken," from his own experiments, as recorded in the communication cited (CHEMICAL NEWS, vol. xl., p. 141). I stated in my letter appearing in CHEMICAL NEWS (vol. xl., p. 155) that "extraordinary care" had been taken by Prof. V. Meyer to make sure he was working with the pure platonic chloride (Pt_2Cl_4 or PtCl_2), and to prove that I made no unfounded statement I will quote the analyses he gives in the *Deut. Chem. Ges. Ber.* for July (xii., p. 1429). He says there "The platonic chloride used by us consisted of a dark olive-green powder, giving on analysis:—

Derivatives of Propionic Acid.—B. Freytag.—A preliminary communication.

Action of Sulphuryl Chloride upon Alcohols.—Dr. P. Behrend.—A reply to P. Claesson.

Dingler's Polytechnisches Journal.
October, 1879.

Chemical Stability of Explosives.—F. Hess.—For practical purposes it is generally sufficient to expose small quantities of the material for eight days in an air-tight vessel to a temperature of 70°. If neither explosion nor the appearance of decomposition products, *e. g.*, hypnitric acid, the substance may be regarded as stable. If the explosive requires to be preserved for years, a more rigorous test is required, which cannot be made intelligible without the accompanying figure.

Apparatus for the production of High Temperatures.—C. A. Paguelin.—Mixtures of combustible gases or vapours with common air are burnt in an especially-constructed receptacle.

Apparatus for determining Solubilities.—H. Kœhler.—Requires the appended figure.

Determination of Iodine in Sea-weeds.—O. Schott.—From the *Zeitschrift f. Anal. Chemie*, 1879, p. 443.—Requires figure.

Examination of Atmospheric Air.—F. Fischer.

Examination and Treatment of Petroleum.—H. Hörler.

Development of Dyeing, Tissue-printing, and Bleaching.—Dr. A. Kilmeyer.—These three papers do not admit of useful abstraction.

Determination of Potassa and Soda in Minerals.—W. Knop and J. Hazard.—The authors dissolve in hydrofluoric acid, evaporate, drench the residue with concentrated sulphuric acid, thus removing the bulk of the silica as silicon fluoride. The sulphuric acid is then evaporated off, the dry residue moistened with five or six drops of concentrated sulphuric acid, heated, drenched with 150 c.c. water, and barium hydrate added till red litmus-paper is turned distinctly blue. The mixture of barium sulphate, silica, alumina, magnesia, and ferric oxide is then filtered off and well washed. The filtrate is evaporated to dryness, adding, when it is concentrated down to about 200 c.c., a few grammes of dry ammonium sesquicarbonate. When perfectly dry the residue of barium and calcium carbonate is extracted successively fifteen times, each time with 20 c.c. water, the liquid being each time filtered through a small filter of 3 to 4 c.m. diameter into a platinum capsule, and evaporated to dryness. The residue is drenched again with 20 c.c. water, the water is decanted through a similar fresh filter, and the solution—after it has deposited a little barium carbonate with some alumina and iron—is collected along with the washings in a fresh platinum capsule. The alkaline carbonate, to which a few more granules of ammonium carbonate have been added, is again dissolved in 20 c.c. water, observing that no residue remains. The liquid is then neutralised with hydrochloric acid, evaporated, the chlorides dried strongly, and the potassium and sodium separated by means of platinum chloride.—*Chem. Central Blatt*.

Moniteur Scientifique, Quasneville.
November, 1879.

On Nitrification.—R. Warington.—From the *Journal of the Chemical Society*.

Manufacture of Carbonate of Potash by Leblanc's Process.—Dr. A. Bugel.—The conditions of this manufacture differ from those of the manufacture of soda, on account of the higher price of the raw material, the greater volatility of potassium compounds at high tem-

peratures, the slight solubility of potassium sulphate concentrated solutions of potassium carbonate, and the non-formation of mother-liquors in the course of the manufacture. It is remarked that if a certain English coal, known as "Ryhope peas," be employed, ferro-cyanide is obtainable up to 1 per cent.

Preparation of Nitro-glycerin.—H. Boutmy.—Not capable of useful abstraction.

On Aurin.—R. S. Dale and C. Schorlemmer.—From the *Journal of the Chemical Society*.

The Phosphates of Lime of Quercy.—M. Rey-Lescure.—An account of the geological position of these deposits, the accompanying fossils, &c.

Origin of Phosphorus in its Various Deposits, and in particular in those of Quercy.—M. Daubrée.—The author holds that phosphorus, notwithstanding its association with fossil bones, is of profound and inorganic origin, its chief source being the eruptive rocks. Meteorites also supply proof of the general distribution of phosphorus in the celestial spaces.

Researches on Amblygonite, and the means of its Utilisation in Agriculture.—M. le Baron P. Thenard.—The substance operated upon contained—

Phosphoric acid	39.00
Alumina	25.66
Ferric oxide	1.71
Lime	1.10
Insoluble matter	10.55
Lithia, potassa, soda, &c.	21.98

100.00

The alumina and the phosphoric acid may be separated from each other, in presence of an excess of sulphuric acid, at 60° (B. ?); but whilst the alumina forms a persulphate insoluble in sulphuric acid, and very slightly soluble in cold water, the phosphoric acid forms a liquid sulpho-conjugated acid. The separation is therefore effected by decanting the sulpho-phosphoric acid and washing the persulphate of alumina.

Active Matter of Malt, or Maltin and Diastase.—M. Dubrunfaut.—The author ascribes the following properties to the diastase of Payen. It exists in malt to the extent of one- to two-thousandths; it liquefies 2000 times its weight of starch; it is free from nitrogen, and saccharifies 2000 times its weight of starch (according to Guerin only 15 times). It has no rotatory power; is soluble in weak alcohol, but insoluble in anhydrous alcohol, and not affected by this liquid. It is not altered on solution in water at 75°, but is modified at 100°. Its maximum of activity is at 75°. Maltin, on the other hand, exists in malt to the extent of 1 part in 100; it liquefies two hundred thousand times its weight of starch; it contains from 0.07 to 0.08 of nitrogen, saccharifies a hundred times weight of starch, and possesses a strong lævo-rotatory power; it dissolves in alcohol at 40 to 50 per cent, but is insoluble in, though modified by, strong alcohol. Its maximum activity is below 50°, and it exists not merely in cereals, but in many potable waters.

Extracts from the Transactions of the Chemical Society of Geneva.—These extracts consist of "Certain Derivatives of the Aromatic Ketones," MM. Friedel, Crafts, Ador, and Billiet; "On Alizarin Blue," C. Graebe; "On Naphtho-picric Acid," M. Bourcart; "On Sulpho-Alizaric Acid," M. Bourcart; "Constitution and Certain Derivatives of Desoxalic Acid," H. Brunner; "Attempts to prepare the Tertiary Bases in which an Atom of Nitrogen is linked by its Three Affinities to Atoms of Carbon differently connected among themselves," H. Bungener; "Chemical and Physical Study of Soils suitable for the Establishment of a Cemetery," L. Lossier; "New Method of the Synthesis of Ketons," G. de Becchi; "Solubility of certain Hydrocarbons and other Aromatic Substances of High Boiling-points in Alcohol and Toluene," G. de Becchi.

Composition of certain Complex Oxides of Cobalt and Nickel.—T. Bayley.—From the *Proceedings of the Royal Irish Academy*.

New Process of Manufacturing Manure with Fresh Urine.—Raoul Brullé.—The author's process, which has been patented, is the absorption of urine by recently baked gypsum.

The Height of "Specialité"—Lourdes-water dialysed by M. Bravais.

Les Mondes, Revue Hebdomadaire des Sciences.
No. 8, October 23, 1879.

Researches on the Constitution of the Ferric Hydrates.—Dr. D. Tommasi.—It appears that all these hydrates may be arranged in two series, respectively isomeric, the red and the yellow. The former are obtained by precipitating a ferric salt by potassa, soda, or ammonia. When calcined they present the phenomenon of incandescence; they dissolve readily, even in the weakest acids. Ferric chloride dissolves them in quantity, and the solution gives a precipitate of ferric hydrate on the addition of potassium sulphate or sulphuric acid. They are dehydrated on boiling with water. The yellow hydrates are obtained by oxidising ferrous hydrate or carbonate or magnetic hydrate. If these hydrates are calcined they do not display incandescence; they are sparingly soluble in acids, whether dilute or concentrated; they are not attacked by ferric chloride; if boiled in water they only lose two molecules of water, retaining the third, even if boiled in a concentrated solution of calcium chloride. It is possible that not merely the ferric salts but also the compounds of chrome and aluminium may exist in these two modifications. If the two ferric chlorides are treated with silver nitrate, the one precipitates all its chlorine, whilst the other gives up 4 atoms at first and the two others afterwards. In a similar manner the blue chromium chloride gives up all its chlorine at once, whilst the green chloride gives up two-thirds at first and the remainder afterwards. There ought to exist two ferric sulphates corresponding to the two chromic sulphates. The two latter are distinguished not merely by their colour, but by the fact that the blue sulphate gives up all its acid to baryta, whilst the green sulphate retains a part.

Report on the Methods of Electric Illumination.—M. Tommasi.—The author points out the disadvantages both of the Bunsen battery and of the dynamo-electric machine as employed for the production of the electric light, and proposes a new battery—a modification of that of Bunsen—which can be worked at 7 centimes per hour per element, whilst a similar intensity, obtained with the dynamo-electric machine, costs 19 centimes. The maintenance of 10 burners with the common Bunsen battery costs 50 centimes per hour; with the magneto-electric machine 25 centimes, and with the Tommasi battery 13 centimes. An equal light produced by means of gas at 30 centimes per cubic metre would cost 42 centimes.

Transactions of the Imperial Academy of Sciences of Vienna.—Mineralogical notes on apophyllite, chabasite, plagioclase, and nephrite.

Die Chemische Industrie.
No. 8, September, 1879.

Apparatus for Concentrating Sulphuric Acid.—J. Stroof.—The author shows that the apparatus of Prentice, whose patent is worked by Messrs. Johnson and Matthey, is far superior to all others. If its performance is taken as 100, that of Faure and Kessler, at equal cost, is 79, that of Desmontis Quennessen and Le Brun 76, and that of the old construction only 38.

Behaviour of the Acids of Nitrogen with Sulphuric Acid.—Dr. G. Lunge.—Already noticed.

Determination of the Value of Superphosphates.—H. Albert and F. Siegfried.—The authors have abandoned the use of ammonium tartrate for dissolving the reverted phosphates, as they find that the action of ammonium citrate at equal temperatures and in equal times is more satisfactory. They find a temperature of 17° to 20° preferable.—*Zeitschrift für Anal. Chemie*.

Annales de Chimie et de Physique,
October, 1879.

Laws of Dispersion of the Dark Thermic Rays and Measurement of their Wave-Lengths.—M. Mouton.—This long and important memoir, of which the first part is here inserted, does not admit of useful abstraction.

Action of Water upon Metallic Sulphides.—MM. de Clermont and Frommel.—The decomposition of metallic sulphides in presence of boiling water is a dissociation of the hydrates of these sulphides effected by heat. The authors prove experimentally that prior to decomposition the sulphides exist as hydrates.

Refrigerating Power of Air at High Pressures.—M. Aimé Witz.—A mathematical paper, not susceptible of useful abstraction.

Researches on the Colouring Matters of Madder.—M. A. Rosenstiehl.—(Fourth and concluding part.)—In this memoir the author describes purpuroxanthin.

Reversion of Superphosphates.—M. H. Joulie.—The author maintains that superphosphates, even when containing much iron and alumina, if prepared with a sufficient quantity of acid, do not undergo a reversion of their assimilable phosphoric acid, but they are apt to remain in a paste-like state. If the dose of acid is smaller, and the action incomplete, the mass dries better, but the assimilable phosphoric acid undergoes reversion. The addition to the superphosphates of calcium carbonate determines immediately the same phenomenon.

Chemical Researches on the Ligneous Papilionaceæ.—MM. P. Fliche and L. Grandea.—Plants belonging to one and the same family and growing on the same soil differ notably in the quantity and distribution of the starch, in the quantity of the ash, and in that of nitrogen. The percentage composition of the ash varies within still wider limits. The nearer genera approach each other the smaller are their chemical differences.

Annalen der Physik und Chemie.
No. 10, 1879.

On Newton's Dust-Rings.—E. Lommel.—An extensive paper, not capable of useful abstraction.

On the Law of Stokes.—E. Lommel.—The author agrees with E. Becquerel, who approves of the conclusion of Lamansky, that the law of Stokes is of universal validity. The experiments of Lamansky, though accurate, merely prove that the mean refrangibility of the light of fluorescence is less than the mean refrangibility of the exciting light.

The Spectrum of Oxygen.—A. Wüllner.—A criticism of the researches of Paalzow, who, in October, 1878, communicated to the Berlin Academy of Sciences the description of an oxygen spectrum, consisting essentially of five lines.

Behaviour of Electricity in Attenuated Gases.—F. Narr.—The author considers it indubitable that the process of eradication cannot be alone explained by the presence of dust, mercurial vapour, or watery vapour.

Electro-magnetic Rotation of the Plane of Polarisation of Light in Gases.—A. Kundt and W. C. Röntgen.—The authors show experimentally that air, oxygen, nitrogen, carbonic oxide, carbonic acid, coal-gas, ethylene, and marsh-gas display a rotation of the plane of polarisation.

sation in the direction of the positive current, like water and carbon disulphide. The magnitude of the rotation differs manifestly for different gases, other conditions being equal.

Theory of the Poles of a Bar-Magnet.—E. Riecke.—An extensive, mathematical paper, not susceptible of useful abridgment.

Constant Batteries of Bunsen, Grove, and Daniell, especially on the Dependence of their Electromotoric Force on the Concentration of the Liquids.—Carl Fromme.—In a Bunsen element the concentration of the nitric acid has a greater influence than in Groves. The stronger the acid the stronger is the current.

Changes of Density in Steel on Hardening.—Carl Fromme.—The hardening of steel is followed by a decrease of its specific gravity.

Journal de Pharmacie et de Chimie.
October, 1879.

Actions of Life without Air; their Influence on the Chemical Phenomena of Respiration.—M. Pasteur.—The author is led to believe that in the animal economy there take place phenomena of the same order as in fermentation. Oxygen acts not only by effecting combustions, but gives to the cellules an activity whence they derive the power of action beyond the influence of free oxygen, in the manner of the ferment-cells. Direct combustions are of little importance.

Chemical Constitution of the Alkaline Amalgams.—M. Berthelot.—Already noticed.

Constitution and Properties of Dialysed Iron.—M. Personne.—The author shows the insolubility of this dialysed iron in the gastric juice.

Constitution of Bodies.—A sketch of some of the recent researches of Mr. N. Lockyer.

New Condensation Hygrometer.—M. Alluard.

The remaining papers in this issue are either purely pharmaceutical or are reprints of matter already noticed from other sources.

Chemiker Zeitung.
No. 40, 1879.

At the meeting of the "International Association against the Pollution of Rivers, Soil, and Air," Professor Vogt, of Bern, declared that according to his experiments the west side of a house was the warmest, upon which followed, in a decreasing series, the east, south, and north. Baron v. Podenils explained his system of dealing with faecal matters. He concentrates by heat, and forces the products of combustion through the mass.

At the meeting of Naturalists and Physicians held at Baden-Baden, Prof. Herman, of Zurich, emphatically denounced the anti-vivisection outcry; Prof. Delffs held a lengthy speech on the behaviour of hydrogen sulphide with the heavy metals; Prof. Schwarz, of Graz, gave an account of the "Coloured Derivatives of Orcin," and Dr. Fithia described a fourth mono-nitrophenol; Dr. Brühl spoke on the exploration of chemical constitution by physical means, a problem which, he maintained, cannot be solved by purely chemical procedures. He proposes the law that the non-saturated compounds possess a molecular refraction apparently too great, the excess above the calculated value being proportional to the number of manifold combinations of adjacent atoms.

Decrease of Volume in the Formation and Decomposition of Aqueous Solutions.—M. Müller-Erbach.—When bodies, solid or liquid, combine with water contraction takes place in almost every instance. In the mutual decomposition of aqueous solutions the contrac-

tion is greater the greater the intensity of the chemical action.—*Chem. Central-Blatt.*

Gay Lussac's Hyponitric Acid.—According to H. Goldschmidt this acid is non-existent. The fumes given off on heating *aqua-regia* consist of a mixture of chlorine and nitrosyl-chloride.—*Chem. Central-Blatt.*

Qualitative Test for Traces of Mercury.—Ed. Teuber.—The material, dry and finely powdered, is mixed with well ignited iron filings and a little red-lead. The mixture is placed in a crucible upon a layer of red-lead, and covered with a layer of iron-filings. The lid of the crucible is so arranged that all the fumes given off on heating the capsule must impinge upon the bottom of a small gold capsule filled with cold water. Traces of mercury down to 0.0001 grm. may thus be detected.—*Oest. Zeitschrift Berg. and Huttenw.*

Limits of the Delectability of Carbonic Oxide.—Dr. W. Hempel.—See *Zeitschrift Anal. Chemie*, xviii., p. 399.

Determination of Commercial Nitrates.—J. de Molins.—The author heats along with potassium hydrate and sulphur. The entire nitrogen of the nitrate is converted into ammonia.—*Monit. Prod. Chim.*

No. 41, 1879.

Distillation of Sulphur with Superheated Steam.—Dr. Gerlach.—The author distils sulphur ores in superheated steam in an apparatus figured in the text, and effects a very considerable increase in the yield as compared with the *calcavone*.

On the Right Use of Chemical Symbols.—Prof. Kraut.—The author urges that except where the constitution of a compound is investigated, the dualistic notation of Berzelius should be used, the atomic weights of the elements being of course corrected.

Congress on Patent Law.—The most important suggestion made is that patents should be granted for chemical products, and not merely as, according to the German law, for the processes by which they are prepared.

New Method for the Detection of Foreign Fats in Butter.—Dr. J. Köttstorfer.—See *Zeitschrift Anal. Chemie*, xviii., p. 431.

Determination of Nitrates in Highly Diluted Solutions.—A. R. Leeds.—For the detection of traces of the oxides of nitrogen by reduction with iron in alkaline solutions the use of caoutchouc stoppers should be avoided. After the introduction of the reducing ingredients water should be distilled out of the apparatus as long as traces of ammonia can be detected in the distillate, and not till then should the substance under examination be introduced into the retort.—*Zeitschrift Anal. Chemie*, xviii., p. 428.

Determination of Caustic Lime in Samples of Crude Burnt Lime.—Dr. H. Bodenbender and Dr. E. Ihlee.—The authors prefer the method of Grandeau. The sample is dissolved in moderately concentrated ammonium nitrate, and the lime thus dissolved is precipitated with ammonium oxalate.—*Zeitschrift Rübens. Ind.*

Detection of Traces of Nitric Acid or Nitrates in Presence of Large Quantities of Nitrous Acid or Nitrites.—A. Piccini.—The substance in question is dissolved in water along with a sufficient quantity of urea, and this liquid is gradually added to a solution of urea and diluted sulphuric acid. When the reaction is over starch-paste and iodide of potassium are added, and if on the introduction of granulated zinc a blue colour appears a nitrate was present.—*Rev. Scient. Ind.*

Examination of Milk.—H. Ohm.—Thirty grms. well-burnt and powdered gypsum are worked up with the milk to a stiff paste at the time of its setting. Milk of 1.030 sp. gr. at 15° C. congealed in 10 hours, with the addition of 25 per cent of water in 2 hours, with 50 per cent in

90 minutes, and with 75 per cent in 40 minutes. Milk skimmed after standing for 24 hours, spec. gr. 1.033, congealed in 4 hours, with 50 per cent water in 1 hour, and with 75 per cent in 30 minutes.—*Arch. Pharm.*, xii., 211.

Detection of Phosphates.—H. Hager.—The substance—food, drinks, excretions, contents of the stomach, &c.—are mixed with basic acetate of lead to remove hydrogen sulphide, and a part of the mixture is then well shaken up with ether in a glass, which is then closed with a cork, in which is inserted a strip of parchment-paper moistened with silver nitrate. If phosphorus is present the paper is quickly turned a lustrous metallic black.—*Pharm. C. H.*, xvii., p. 327.

MISCELLANEOUS.

The Physical Society.—In addition to the papers announced in our last issue to be read at the meeting to-morrow, Dr. Guthrie, F.R.S., will read a paper on "Prof. Pisani's experiments on the Incandescence of Electrodes."

Society of Arts.—The Programme of this Society for its 126th Session has just been issued. It gives a list of the Papers and Lectures for the Session, so far as they have been arranged. The following are the papers to be read at the evening meetings previous to Christmas:—November 26—"Suggestions for Dealing with the Sewage of London," by Major-General H. Y. D. Scott, C.B., F.R.S. December 3—"Apprenticeship: Scientific and Unscientific," by Sylvanus P. Thompson, D.Sc., Professor of Applied Physics at University College, Bristol. December 10—"Art Vestiges in Afghanistan; the Results of Some Recent Explorations in the Jellalabad Valley," by William Simpson. December 17—"The Panama Canal," by Captain Bedford Pim, R.N., M.P. The dates of the Papers after Christmas are not announced, but the following are among the subjects to be treated:—"Domestic Poisons," by Henry Carr; "Gas Furnaces and Kilns for Burning Pottery," by Herbert Guthrie, C.E.; "The Utilisation of Slag," by Charles Wood; "Art in Japan," by C. Pfouder; "The Trade and Commerce of the Yenisei," by Henry Seebohm; "Modern Autographic Printing Processes," by Thomas Bolas, F.C.S.; "The History of the Art of Bookbinding," by Henry B. Wheatley, F.S.A.; "Art Ironwork," by J. W. Singer; "The History of Musical Pitch," by A. J. Ellis, F.R.S.; "The Recent History of Explosive Agents," by Professor Abel, C.B., F.R.S.; "Ireland and its Resources," by C. G. W. Lock; "The Future of Epping Forest," by William Paul, F.L.S. Three courses of "Cantor Lectures" are to be given. The First Course is by Dr. Charles Graham, F.C.S., F.I.C. Professor of Chemical Technology at University College, London, on "The Chemistry of Bread and Bread-making." The Second on the "Manufacture of India-rubber and Gutta-percha," by Thomas Bolas, F.C.S. The Third by R. W. Edis, F.S.A., "Art Decoration and Furniture." The first meeting of the Session will be held on the 19th inst., when the Opening Address will be delivered by Lord Alfred S. Churchill, Chairman of the Council.

NOTES AND QUERIES.

Chlorophyll.—Can anyone say where chlorophyll may be obtained in quantity?—*CHLOR.*

TO CORRESPONDENTS.

A. McDougall.—The process has not yet been published. No particulars are known.

"Address to Students," *C.N.*, Sept. 12, 1879.—The Editor wishes to remove an impression which prevails respecting the authority for the case given of the student who carried off the B.A. prize as well as Honours in Chemistry by dint of an excellent memory. This case was not given on the authority of Professor Huxley, but on that of the writer of the article.

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AND
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THE CHEMICAL NEWS.

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THE PERIODIC LAW OF THE CHEMICAL ELEMENTS.

By D. MENDELBERG.

ALTHOUGH seven years have passed since these thoughts absorbed any attention; although other occupations have drawn my attention from the problem of the elements, which was always getting nearer solution; in short, although I might wish to put this question otherwise than I did seven years ago, still I keep to the same firm conviction that I formerly had on the importance and value of the theorems on which my memoir is based; and this is why I am so gratified to see my ideas exposed to the appreciation of the learned. Several occurrences have aided to make some of the logical consequences of the periodic law popular.

1st. The law I announced has been considered as a repetition in another form of what has already been said by others. The article in the *Berichte der Deutschen Chemischen Gesellschaft*, 1874, p. 348, treats only of the question of priority. It is now certain that the periodic law offers consequences that the old system had scarcely ventured to foresee. Formerly it was only a grouping, a scheme, a subordination to a given fact; while the periodic law furnishes the facts, and tends to strengthen the philosophical question, which brings to light the mysterious nature of the elements. This tendency is of the same category as Froust's law, with the essential difference that Froust's law is arithmetical, and that the periodic law exhausts itself in connecting the mechanical and philosophical laws which form the character and glory of the exact sciences. It proclaims loudly that the nature of the elements depends above all on their mass, and it considers this function as periodic. The formula of the law might be changed; a greater appreciation of this function will be found; but, I believe that the original idea of the periodic law will remain, because it is opposed to the primitive notion that the nature of elements depends on unknown causes, and not on their mass. In 1872 to 1877 Gustavson and Palitzine showed that even the coefficients of simple substitution depend on the atomic weight of elements, in the sense which is in accordance with the periodic law.

2ndly, The periodic law requires changes in the atomic weights of many elements yet incompletely examined; for example, indium, cerium, yttrium, erbium, didymium, &c. Before the existence of this law there was no reason for doubting the generally accepted atomic weights of

* Considerable attention having been drawn to M. Mendeleef's memoir "On the Periodic Law of the Chemical Elements," in consequence of the discovery of the elements gallium and scandium being apparently identical with his two predicted elements ekaluminium and ekaboron, it has been thought desirable to reproduce the entire article in the CHEMICAL NEWS. The translation is from the *Moniteur Scientifique*, July, 1879, to which periodical it has been communicated by M. Mendeleef, who prefaces the memoir by letter. This letter forms the subject of the present article; the memoir will be commenced next week.—Ed. C.N.

† Touching the compressibility, the dilation, and the resistance of gases, the temperature of the upper layers of the atmosphere, aerostatics, researches on the origin of petroleum, &c.

these elements. At present, since the researches of Ramsberg, Roscoe, Clève, Høglund, Hildebrand, and others, have led to the same conclusion as that which is derived from the periodic law, it becomes incontestable that it shows us the truth, which is in fact its support.

3rdly. The periodic law has given the first chance of predicting not only unknown elements, but also of determining the chemical and physical properties of simple bodies still to be discovered, and those of their compounds. The discovery of gallium by M. Lecoq de Boisbaudran, whom I have now the honour of counting as a friend, may be considered as the inauguration of the periodic law, and reckoned amongst the brilliant pages in the annals of science. The properties announced in the *Comptes Rendus*, Nov. 22, 1875, have been confirmed by after study. It will be enough to mention four—the formation of gallium alum, the equivalent of the oxide, the specific gravity of the metal (5.9), and the atomic weight of the element.

It must be admitted that before the periodic law there was no possibility of a similar prediction. It is here to the point to direct attention to the fact that the fusing-point of gallium is so low that it melts at the heat of the hand. This property might be considered as foreseen, but that is not the case; it suffices to look at the following series:—

Mg	Al	Si	P	S	Cl
Zn	Ga	—	As	Se	Br
Cd	In	Sn	Sb	Te	I

It is evident that in the group Mg, Zn, Cd, the most refractory element (Mg) has the lowest atomic weight; but in the groups commencing with S and Cl the most difficultly fusible bodies are the heaviest. In a transitory group, as Al, Ga, In, it is necessary to notice an intermediate fact, the extremes, that is, the heaviest (In) and the lightest (Al) should be less fusible than the middle one, as it is in reality. I would further remark that a property such as the fusion-point is characterised chiefly by the molecular weight and not by the atomic weight. If we had a variation of solid sulphur, not as S^{VI} (or perhaps heavier, as S^8) but S^{II} , which form it acquires at 800° , its boiling- and fusion-point would have been much lower. In the same way, ozone, O^{III} , will condense much more easily than oxygen, O^{II} . Experiments made in our laboratory confirm this fact.

The three circumstances mentioned have obliged chemists to direct their attention, distracted by the brilliant material acquisitions which distinguish our epoch, towards the periodic law. I ought now myself to complete what is still wanting on this subject, but for the time being I am occupied with other matters, and I should leave the care of developing this question to the future and to new forces, which will I hope endeavour to bring as the first fruits of the periodic law a new philosophical order, in fixing it by pillars strengthened by new experiments, so as to give greater stability to the edifice already begun. I will add but three brief observations:—

1st. The best way of drawing up the table of elements, so as to show the periodic relations, will, I think, be:—

TYPICAL ELEMENTS.

I.	II.	III.	IV.	V.	VI.	VII.
H						
Li	Be	B	C	N	O	F
Na						

EVEN ELEMENTS.

I.	II.	III.	IV.	V.	VI.	VII.
K	Ca	—	Tl	V	Cr	Mn
Rb	Sr	Yt	Zr	Nb	Mo	—
Cs	Ba	La	Ce	—	—	—
—	—	Er	Di (?)	Ta	W	—
—	—	—	Th	—	U	—

ODD ELEMENTS.

VIII.			I.	II.	III.	IV.	V.	VI.	VII.
Fe	Co	Ni	Cu	Mg	Al	Si	P	S	Cl
Ru	Rh	Pd	Ag	Zn	Ga	In	As	Se	Br
Os	Ir	Pt	Au	Hg	Tl	Pb	Bi		I

The Roman figures show the groups or the forms of combination.

2nd. With regard to new elements recently discovered, I think it necessary to keep silence. Of late years metals have been born and have died, such as Davyum, Mosandrium, &c.; this shows the necessity of caution. It is only necessary to mention Marignac's ytterbium (*Archives des Sciences Physiques et Naturelles*, 1878, Nov. 15, No. 251), because the name of the investigator is itself sufficient guarantee. But, after announcement of the series of new metals of Delafontaine, he himself asked that someone else should make fresh researches in such an inaccessible region as gadolinite. One can still say the same of oxide of didymium; it should be re-examined. The absorption-bands of solutions and the equivalents of oxides are not sufficiently sure to show the individuality of elements; because, in different degrees of oxidation and in salts of different basicity, the absorption-bands and the equivalents of oxides can be heterogeneous for the same elements, as, for instance, in cerium, uranium, iron, chromium, &c.

3rd. I should like to fix the attention of chemists on the three principles which are demonstrated in the last chapter, and also on the consequences which are adduced from their strict application to organic compounds (above all not saturated): by their aid we do without hypotheses, we explain cases of isomerism, and we obtain new consequences as yet unexplained.

D. MENDELEEF.

(To be continued.)

THE ALLEGED DECOMPOSITION OF CHORINE.

DR. H. ENDEMANN, who has written an article in the *Journal of the American Chemical Society* entitled "A Review of the Latest Investigations on the Dissociation of the Elements at High Temperatures," publishes the following letter from Prof. Meyer:—

"Zurich, d 28 Sept., 1879.

"SEHR GEHRETER HERR,—In Erwiderung auf Ihr geehrtes Schreiben vom 12 Sept., theile ich Ihnen mit, dass die Mittheilungen der englischen Journale ueber meine Arbeiten welche ueber Gewinnung von Sauerstoff aus Chlor berichten, ohne mein Wissen und gegen meinen Willen veröffentlicht sind, und das diese Mittheilungen in wesentlichen Punkten voellig inkorrekt sind.—Hochachtungsvoll,

"Prof. VICTOR MEYER."

[TRANSLATION.]

"Zurich, September 28, 1879.

"DEAR SIR,—In reply to your favour of September 12th, I would state that the communications in the English journals concerning my investigations which report the separation of oxygen from chlorine, have been published without my knowledge and against my wishes, and that the said communications are entirely incorrect in essential particulars.—Respectfully,

"Prof. VICTOR MEYER."

BLOWPIPE ANALYSIS.

THE increasing appreciation of the value of Blowpipe Analysis in England is shown by the recent publication of the translation of another German work on the subject by one of our first scientific publishers, Messrs. Macmillan.

We shall take an early opportunity of reviewing this little work by Herr Landauer, of Brunswick, who lately visited England at the head of a sanitary commission sent here by that State, but meantime have the pleasure

to offer our readers extracts from a letter to Colonel Ross, whose contributions on the subject have so often appeared in this journal. The writer of this letter, whose name we are not at present at liberty to communicate, is obviously one well acquainted with the subject, but we may mention that he is a credit to a body of scientific officers who are admitted throughout Europe to be a credit to this country, the Corps of Royal Engineers.

"Point de Galle, Ceylon,

"Sept. 13th, 1879.

"DEAR SIR,—I have to acknowledge your letter, with thanks for the information it contained. I long since saw notices of your new system of Blowpipe Analysis, one stating that your work was the best English authority on the subject. It was only recently that I got out "Griffin's Handicraft," and going over the list of your apparatus I saw that your method must be something very different from the old, so I ordered out a special box both for the blowpipe and separately for the wet way. Your book came to hand afterwards. There is no question that your method is greatly superior to Plattner's, which I think may be considered out of date, and that it is a powerful and logical mode of chemical analysis. . . . I was greatly pleased both with boric and phosphoric acid as vehicles of decomposition, and especially so with the lime borate balls. . . . Your book I found extremely interesting, as one written by an enthusiastic master of his subject can hardly fail to be. From your going to publish an epitome, it is evident that you think with myself that Spon's edition is rather too bulky. . . .

The chapter on flame, apparatus, reagents, manipulation can hardly be better, being very intelligible and practical. The apparatus is a great advance on the old kinds. . . . The chapter on colour seems to me to be not in the right place, but I have not as yet made any particular use of analysis by colour. . . . I generally go when in search of information to the next chapter, where the ordinary metals and bases are dealt with. This is very clear throughout, and easily worked from. . . . The last half of your book appeals to three classes of investigators. It might, therefore, form usefully three divisions—one for mineralogists, who would be chiefly geologists and engineers; the next to relate to metals intended for the guidance of chemists and manufacturers; and the third for the use of farmers. This division would be very convenient, because one generally employs the blowpipe according to locality, and oftener for some specific branch than for mere study of the multitudinous specimens, seldom found in any number together out of a cabinet or museum. As to the chapter on farming, it almost seems necessary to follow the order of some textbook: "Church's Laboratory Guide" (Macmillan) is one of the best on agricultural chemistry that I know. What is required would be to show *seriatim* how all that he accomplishes in the wet way can be attained by the blowpipe, including the physical properties of the soil. The best way of in-uring this would be to go right through the book, altering it in brief to blowpipe methods. At present one has to do this each time, wading backwards and forwards through descriptions in pyrology intended for the general student till a result of some kind is obtained. . . . A working chapter on the blowpipe applied to agricultural chemistry would supply a great want. . . . I am more familiar with the wet processes, and as Church lays down very complete instructions can depend upon what I would obtain by this method. But there is no reason except this, I think, for considering the wet way preferable; in fact, the blowpipe system now rests upon exactly the same numerical basis the other does, and the apparatus is much more compact, manageable, and cheap.

"There are, however, processes, such as the volumetric analysis of water, or the analysis of water generally, which do not appear to come within the scope of the blowpipe.—Yours very faithfully,

"A. F. F."

ON THE VAPOUR-DENSITIES OF
PEROXIDE OF NITROGEN, FORMIC ACID,
ACETIC ACID, AND PERCHLORIDE OF
PHOSPHORUS.

By J. WILLARD GIBBS.

(Continued from p. 224.)

Peroxide of Nitrogen.—If we take the constants of the equation for this substance from the paper already cited,* we have—

$$\log \frac{15.89(D-1.589)}{(3.178-D)^2} = \frac{3118.6}{t_c + 273} + \log p - 12.451, \quad (10)$$

t_c denoting the temperature on the centigrade scale. The numbers 3.178 and 1.589 represent the theoretical densities of N_2O_4 and NO_2 respectively. The two other constants were determined by the experiments of Deville and Troost.

The results of these and other experiments at atmospheric pressure, all made by Dumas's method, are exhibited in Table I. The first three columns give the temperature (centigrade), the pressure (in millimetres of mercury),† and the density calculated from the temperature and pressure by equation (10). The subsequent columns give the densities observed by different authorities, and the excess of the observed over the calculated densities. In the first column of observed densities we have one observation by Mitscherlich‡ (at 100.25°) and five by R. Müller.§ The three remaining columns contain each the results of a series of experiments by Deville and Troost.|| In each series the experiments were made with increasing temperatures, and with the same vessel without refilling. It should be observed that the results of the three series are not regarded by their distinguished authors as of equal weight. It is expressly stated that the numbers in the two earlier series, and especially in the first, may be less exact. The last series agrees very closely with the formula. It was from this that the constants of the formula were determined. The experiments of series I. and II., and those of Mitscherlich and Müller, give somewhat larger values, with a single exception, as is best seen in the columns which give the excess of the observed density. The differences between the different columns are far too regular to be attributed to the accidental errors of the individual observations, except in the case of the experiment at 151.8°, where some accident has evidently occurred either in the experiment itself or in the reduction of the result. Setting this observation aside, we must look for some constant cause for the other discrepancies between the different series.

We can hardly attribute these discrepancies to difference in the material employed, or to air or other foreign substance imperfectly expelled from the flask. For impurities which increase the density would make the divergence between the different series greatest when the densities are the least, whereas the divergences seem to vanish as the density approaches the limiting value. (A similar objection would apply to the supposition of any error in the determination of the weight of the flask when filled with air alone.) But if we should attribute the divergen-

ces to an impurity which diminishes the density (as air), we should be driven to the conclusion that the first series of Deville and Troost gives the most correct results, and that all the best attested numbers at temperatures below 90° are considerably in the wrong. It does not seem possible to account for these discrepancies by any causes which would apply to cases of normal or constant density. They are illustrations of the general fact that when the density varies rapidly with the temperature, determinations of density for the same temperature and pressure by different observers, or different determinations by the same observer, exhibit discordances which are entirely of a different order of magnitude from those which occur with substances of normal or constant densities, or which occur with the same substance at temperatures at which the density approaches a constant value. In some cases such results may be accounted for by carelessness on the part of the observers, not controlled by a comparison of the result with a value already known. But such an explanation is inadequate to explain the general fact, and evidently inadmissible in the present case.

It is probable that these discrepancies are in part attributable to a circumstance which has been noticed by M. Wurtz, in his account of his experiments upon the vapour-density of bromhydrate of amylene, in the following words:—"Le temps pendant lequel la vapeur est maintenue à la température où l'on détermine la densité n'est pas sans influence sur les nombres obtenus. C'est ce qui résulte des deux expériences faites à 225 degrés avec des produits identiques. Dans la première, la vapeur a été portée rapidement à 225 degrés. Dans la seconde elle a été maintenue pendant dix minutes à cette température. On voit que les nombres trouvés pour les densités ont été fort différents. [The numbers were 4.69 and 3.68 respectively.] Ce résultat ne doit point surprendre si l'on considère que le phénomène de décomposition de la vapeur doit absorber de la chaleur, et que les quantités de chaleur nécessaires pour produire et la dilatation et la décomposition ne sauraient être fournies instantanément."

It is not difficult to form an estimate of the quantities of heat which come into play in such cases. With respect to peroxide of nitrogen, it was estimated in the paper already cited that the heat absorbed in the conversion of a unit of N_2O_4 into NO_2 under constant pressure is represented by $7181 a_2$. (The heat is supposed to be measured in units of mechanical work.) Now the external work done by the conversion of a unit of N_2O_4 into NO_2 under constant pressure is $a_2 t$. Therefore, the ratio of the heat absorbed to the external work done by the conversion of N_2O_4 into NO_2 is $7181 + t$, or 23 at the temperature of 40° C. Let us next consider how much more rapidly this vapour expands with increase of temperature at constant pressure than air. From the necessary relation—

$$v = \frac{k m t}{p D}$$

where m denotes the weight of the vapour, and k a constant, we obtain—

$$\left(\frac{dv}{dt}\right)_p = \frac{v}{t} - \frac{v}{D} \left(\frac{dD}{dt}\right)_p,$$

where the suffix p indicates that the differential coefficients are for constant pressure. The last term of this expression evidently denotes the part of the expansion which is due to the conversion of N_2O_4 into NO_2 , and the preceding term the expansion which would take place if there were no such conversion, and which is identical with the expansion of the same volume of air under the same circumstances. The ratio of the two terms is—

$$-\frac{t}{D} \left(\frac{dD}{dt}\right)_p,$$

the numerical value of which for the temperature of 40° is 2.42, as may be found by differentiating equation (10), or,

* See equation (336) on page 339, *loc. cit.*; also the following equations in which the density is given in terms of the temperature and pressure. In comparing these equations, it must be observed that in (336) the pressures are measured in atmospheres, but in this paper in millimetres of mercury.

† 760 m.m. has been assumed as the pressure of the atmosphere in all cases in which the precise pressure is not recorded in the published account of the experiments. The figures inserted in the columns of pressures are in such cases enclosed in parentheses. The same course has been followed in the subsequent tables. With respect to the principal series of observations by Deville and Troost (series III.), it is stated that the barometer varied between 747 and 764 millimetres. A difference of 13 millimetres in the pressure would in no case cause a difference of 0.005 in the calculated densities. In this series, therefore, the errors due to this circumstance are not very serious.

‡ *Ann. Chim. Phys.*, vol. xxix. (1833), p. 220.

§ *Liv. Ann.*, vol. cxxii. (1862), p. 15.

|| *Comptes Rendus*, vol. lxi. (1867), p. 237.

* *Comptes Rendus*, t. lx, p. 730.

with less precision, from the numbers in the third column of Table I. Let us now suppose that equal volumes of peroxide of nitrogen and of air at the temperature of 40° and the pressure of one atmosphere receive equal infinitesimal increments of temperature under constant pressure. The heat absorbed by the peroxide of nitrogen on account of the conversion of N_2O_4 into NO_2 is 23 times the external work due to the same cause, and this work is 2.42 times the external work done by the expansion of the air. But the heat absorbed by the air in expanding under constant pressure is well known to be 3.5 times the work done. Therefore the heat absorbed on account of the conversion of N_2O_4 into NO_2 is $(23 \times 2.42 \div 3.5) = 15.9$ times the heat absorbed by the air. To obtain the whole heat absorbed by the vapour we must add that which would be required if no conversion took place. At 40° the vapour of peroxide of nitrogen contains about 54 molecules of N_2O_4 to 46 of NO_2 , as may easily be calculated from its density. The specific heat for constant pressure of a mixture in such proportions of gases of such molecular formulæ, if no chemical action could take place, would be about twice that of the same volume of air. Adding this to the heat absorbed by the chemical action, we obtain the final result,—that at 40° and the pressure of the atmosphere the specific heat of peroxide of nitrogen at constant pressure is about eighteen times that of the same volume of air.*

But the greater amount of heat which is required to bring the vapour to the desired temperature is only one factor in the increased liability to error in cases of this kind. The expansion of peroxide of nitrogen for increase of temperature under constant pressure at 40° is 3.42 times that of air. If then, in a determination of density, the vapour fails to reach the temperature of the bath, the error due to the difference of the temperature of the vapour and the bath will be 3.42 times as great as would be caused by the same difference of temperatures in the case of any vapour or gas having a constant density. When we consider that we are liable not only to the same, but to a much greater difference of temperatures in a case like that of peroxide of nitrogen, when the exposure to the heat is of the same duration, it is evident that the common test of the exactness of a process for the determination of vapour-densities, by applying it to a case in which the density is nearly constant, is entirely insufficient.

That the experiments of the III. series of Deville and Troost give numbers so regular and so much lower than the other experiments is probably to be attributed in part to the length of the time of exposure to the heat of the experiment, which was half an hour in this series: for the other series the time is not given.

Another point should be considered in this connection. During the heating of the vapour in the bath it is not material whether the flask is open or closed. This will appear if we compare the values of—

$$\left(\frac{dD}{dt}\right)_p \quad \text{and} \quad \left(\frac{dD}{dt}\right)_v,$$

the differential coefficients of the density with respect to the temperature on the suppositions, respectively, of constant pressure and of constant volume. For 40° we have—

$$\left(\frac{dD}{dt}\right)_p = 0.0189, \quad \left(\frac{dD}{dt}\right)_v = 0.0163,$$

the first number being obtained immediately from equation (10) by differentiation after substitution of—

$$\frac{hmt}{vD} \text{ for } p.$$

The ratio of these numbers evidently gives the proportion in which the chemical change takes place under the two

suppositions. This shows that only about six-sevenths of the heat required for the chemical change can be supplied before opening the flask, and the remainder of this heat, as well as that required for expansion, must be supplied after the opening. The errors due to this source may evidently be diminished by diminishing the intervals of temperature between the successive experiments in a series of this kind, and also by diminishing the opening made in the flask, which increases the time for which the flask may be left open without danger of the entrance of air. In series III. of experiments by Deville and Troost the intervals of temperature did not exceed ten degrees (except after the density had nearly reached its limiting value), and the neck of the flask was drawn out into a very fine tube.

In Table II., which relates to experiments on the same substance at pressures less than that of the atmosphere, the principal series is that of Naumann,* which commences a few degrees below the lowest temperatures of Deville and Troost, and extends to $-6^{\circ} C.$, the pressures varying from 301 to 84 millimetres. These experiments were made by the method of Gay-Lussac. The numbers in the column of observed densities have been re-calculated from the more immediate results of the experiments, and are not in all cases identical with those given in Professor Naumann's paper. Every case of difference is marked with brackets. Instead of the numbers [2.66], [2.62], [2.85], [2.94], Naumann's paper has 2.57, 2.65, 2.84, 3.01, respectively. In some cases the temperatures and pressures of two experiments are so nearly the same that it would be allowable to average the results, at least in the column of excess of observed density. In such cases the numbers in this column have been united by a brace. The greatest difference between the observed and calculated densities is 0.16, which occurs at the least pressure, 84 millimetres. In this experiment the weight of the substance employed is also less than in any other experiment. Under such circumstances the liability to error is of course greatly increased. The average difference between the observed and calculated densities is 0.063. Since these

TABLE II.—PEROXIDE OF NITROGEN.

Experiments at less than Atmospheric Pressure.

PLAYFAIR AND WANKLYN—TROOST—NAUMANN.

Temp.	Pres- sure.	Density calc. by eq. (10).	Density observed.			Excess of obs. density.		
			P. & W.	T.	N.	P. & W.	T.	N.
97.5 (301)		1.631	1.783			+ .152		
27	35	1.90		1.6			— .30	
27	16	1.77		1.59			— .18	
24.5 (323)		2.524	2.52			— .004		
22.5	136.5	2.34		2.35				+ .01
22.5	191	2.26		2.28				+ .02
21.5	161	2.41		2.38				— .03
20.8	153.5	2.41		2.46				+ .05
20	301	2.59		2.70				+ .11
18.5	136	2.43		2.45				+ .02
18	279	2.61		2.71				+ .10
17.5	172	2.51		2.52				+ .01
16.8	172	2.53		2.55				+ .02
16.5	224	2.59		[2.66]				+ .07
16	228.5	2.61		[2.62]				+ .01
14.5	175	2.58		2.63				+ .05
11.3 (199)		2.620	2.645			+ .025		
11	190	2.66		2.76				+ .10
10.5	163	2.64		2.73				+ .09
4.2 (129)		2.710	2.588			— .122		
4	172.5	2.77		2.85				+ .08
2.5	145	2.76		[2.85]				+ .09
1	138	2.78		2.84				+ .06
—1	153	2.83		2.87				+ .04
—3	84	2.76		2.92				+ .16
—5	123	2.85		2.92				+ .07
—6	125.5	2.87		[2.94]				+ .07

* Similar calculations from less precise data for the bromhydrate of amylenes at 225° seem to indicate a specific heat as much as forty times as great as that of the same volume of air.

* Ber. der Deutsch. Chem. Gesell., Jahrgang xl. (1893), S. 225.

differences are almost universally positive, and increase as the temperature diminishes, it is evident that they might be considerably diminished by slight changes in the constants of equation (10) without seriously impairing the agreement of that equation with the experiments of Deville and Troost. But it has not seemed necessary to re-calculate the formula, which, in its present form, will at least illustrate the degree of accuracy with which densities at low pressures and at temperatures below the boiling-point of the liquid may be derived from experiments at atmospheric pressure above the boiling-point. Moreover, the excess of observed density may be due in part to a circumstance mentioned by Professor Naumann, that the chemical action between the vapour and the mercury diminished the volume of the vapour, and thus increased the numbers obtained for the density.

The same table includes two experiments of Troost,* by Dumas's method, but at the very low pressures of 35 m.m. and 16 m.m. In such experiments we cannot expect a close agreement with the formula, for the same error in the determination of the weight of the vapour, which would make a difference of 0.01 in the density in experiments at atmospheric pressure, would make a difference of 0.21 or 0.47 in the circumstances of these experiments. In fact, the numbers obtained differ considerably from those demanded by the formula.

There remain four experiments by Playfair and Wanklyn,† in which Dumas's method was varied by diluting the vapour with nitrogen. The numbers in the column of pressures represent the total pressure diminished by the pressure which the nitrogen alone would have exerted. They are not quite accurate, since the data given to the memoir cited only enable us to determine the ratios of the total and the partial pressures. The numbers here given are obtained by setting the total pressure, which was that of the atmosphere at the time of the experiment, equal to 760 m.m. The effect of this inaccuracy upon the calculated densities would be small. Two of these observations agree closely with the formula; and two show considerable divergence, but in opposite directions, and these are the two in which the quantities of peroxide of nitrogen were the smallest. The differences appear to be attributable rather to the difficulty of a precise determination of the quantities of nitrogen and of vapour than to any effect of the one upon the other.

Special interest attaches to experiments at the same or nearly the same temperature, but at different pressures. For with experiments at the same temperature, the constants of the formula which are determined by observation are reduced to one, so that the verification of the formula by experiment cannot possibly be regarded as a case of interpolation. It is not necessary that the temperatures should be exactly the same, for it will be conceded that the formula represents the actual function well enough to answer for adjusting slight differences of temperature; but it is necessary that the range of pressures should be considerable in order that the differences of density should be large in proportion to the probable errors of observation. But the pressures must not be so low that accurate determinations become impossible.

In the experiments of Naumann we see some fair correspondences with the formula in respect to the influence of pressure, especially in the first four experiments of the list, where, if we average the results of the third and fourth experiments, as is evidently allowable, the observed values follow very closely the fluctuations of the calculated, extending from 2.26 to 2.41. In other cases the agreement is less satisfactory. The circumstance that the experiments at the two highest pressures (301 and 279 m.m.) give results exceeding the two calculated values considerably more than any other experiments at adjacent temperatures may seem to indicate that the densities increase with the pressure more rapidly than the formula allows; but the differences are not too large to be ascribed

to errors of observation, and the experiment at the lowest pressure (84 m.m.) also shows a large excess of observed density.

A much more critical test may be found in the comparison of Naumann's experiments with those of Deville and Troost, notwithstanding the interval of about 4° of temperature. The formula requires that a diminution of pressure from 760 to 101 millimetres shall reduce the density from 2.676 at 26.7° to 2.26 at 22.5, notwithstanding the effect of the change of temperature. Experiment gives a reduction of density from 2.65 to 2.28, which is about one-ninth less. This is, it will be observed, a deviation from the formula in the opposite direction from that which the experiments of Naumann alone, or a comparison of the experiments of Troost with those of Deville and Troost, seem to indicate. The experiment here compared with Naumann's belongs to series III. of Deville and Troost. If, instead of this experiment, we should take an average of the experiments at lowest temperatures in series II. and III., the agreement with the formula with respect to the effect of change of pressure would be almost perfect.

Formic Acid.—In Table III. the determinations of

TABLE III.—FORMIC ACID.

Experiments of Bineau.

Temperature.	Pressure.	Density calculated by eq. (11).	Density observed.	Excess of observed density.
216.0	690	1.60	1.61	+ .01
184.0	750	1.64	1.68	+ .04
125.5	687	2.03	2.05	+ .02
125.5	645	2.02	2.03	+ .01
124.5	670	2.04	2.06	+ .02
124.5	640	2.03	2.04	+ .01
118.0	655	2.13	(2.14)	(+ .01)
118.0	650	2.13	2.13	.00
117.5	688	2.15	2.13	— .02
115.5	649	2.17	2.20	+ .03
115.5	640	2.16	2.16	.00
115	655	2.18	(2.13)	(— .05)
111.5	690	2.25	2.22	— .03
111.5	690	2.25	2.25	.00
111	608	2.22	(2.13)	(— .09)
108	[687]	2.30	2.31	+ .01
105.0	691	2.35	2.35	.00
105.0	650	2.34	2.33	— .01
105.0	630	2.33	2.32	— .01
101.0	693	2.42	2.44	+ .02
101.0	650	2.40	2.41	+ .01
99.5	690	2.44	2.52	+ .08
99.5	684	2.44	2.49	+ .05
99.5	676	2.44	2.46	+ .02
99.5	662	2.43	2.44	+ .01
99.5	641	2.42	2.42	.00
99.5	619	2.41	2.41	.00
99.5	602	2.41	2.40	— .01
99.5	557	2.39	2.34	— .05
34.4	28.94	2.82	2.77	— .05
31.5	3.04	2.40	2.60	+ .20
30.5	8.83	2.67	2.69	+ .02
30.0	18.28	2.81	2.76	— .05
29.0	27.40	2.88	2.83	— .05
24.5	17.39	2.88	2.86	— .02
22.0	25.17	2.95	3.05	+ .10*
20.0	16.67	2.93	2.94	+ .01
20.0	7.99	2.84	2.85	+ .01
20.0	2.72	2.64	2.80	+ .16
18.5	23.53	2.98	3.23	+ .25*
16.0	15.97	2.97	3.13	+ .16*
15.5	2.61	2.72	2.86	+ .14
15.0	7.60	2.90	2.93	+ .03
12.5	15.20	3.00	3.14	+ .14*
11.0	7.26	2.95	3.02	+ .07
10.5	14.09	3.01	3.23	+ .22*

* *Comptes Rendus*, t. lxxvi. (1878), p. 1395.

† *Trans. Roy. Soc. Edin.*, vol. xxii. (1861), p. 463.

Bineau are compared with the densities calculated by the formula—

$$\log \frac{1.589(D - 1.589)}{(3.178 - D)^2} = \frac{3800}{t_c + 273} + \log p - 12.641 \quad (11)$$

The observed densities are taken from the eighteenth volume of the third series of the *Annales de Chimie et de Physique* (1846), except in three cases, distinguished by parentheses, which are earlier determinations published in the nineteenth volume of the *Comptes Rendus* (1844). It may be added that the pressure (687) for the experiment at 108° is taken from Erdmann's *Journal für Praktische Chemie* (vol. xl., p. 44), the impression being imperfect in the *Annales*, in the copies to which the author has been able to refer, where the figures look much like 637. (The pressure 637 would make the calculated density 2.28.)

In the column which gives the excess of observed densities, the effect of nearness to the state of saturation is often very marked. Such cases are distinguished by an asterisk. The temperature of 99.5° is below the boiling-point of formic acid, and the higher pressures employed at this temperature cannot be far from the pressure of saturated vapour. With respect to lower temperatures, we have the statement of Bineau that the pressure of saturated vapour is about 19 m.m. at 13°, 20.5 m.m. at 15°, 33.5 m.m. at 22°, and 53.5 m.m. at 32°. By interpolation between the *logarithms* of these pressures (in a single case, by *extrapolation*), we obtain the following result:—

Temperature	10.5	12.5	16	18.5	22
Pressure of sat. vapour	16.6	18.5	22	26.2	33.5
Pressure of experiment	14.69	15.20	15.97	23.53	25.17

Whether the large excess of observed density in these cases represents a property of the vapour, or an incipient condensation on the walls of the vessel which contains it, as has been supposed by eminent physicists in similar cases, we need not here discuss.

If we reject these cases of nearly saturated vapour, as well as the three earlier determinations, there remain twenty-five experiments at pressures somewhat less than one atmosphere in which the maximum difference between the observed and calculated densities is 0.05, and the average difference 0.016; nine experiments at pressures ranging from 29 m.m. to 7 m.m., in which the maximum difference is 0.07 and the average 0.035; and three experiments at pressures of about 3 m.m., in which the average difference is 0.17. The extraordinary precision of the determinations at low pressures is doubtless due to the large scale on which the experiments were conducted. All the experiments at temperatures below 99° were made with a globe of the capacity of 5½ litres, with a stem of suitable length to hold the barometric column.

The agreement is certainly as good as could be desired, and shows the accuracy of which the method of observation is capable. But in no part of the thermometric scale do we find so great a range of pressures as might be desired, without using pressures too low for accurate results, or observations which are to be rejected for other reasons.—*American Journal of Science*.

PECULIAR REACTION OF AMMONIA ON BRASS.

By JOHN Y. McLELLAN.

WHILE experimenting on the action of liquor ammonia on various metals and alloys with a view to determine the most suitable for the contraction of a certain part in an ammonia plant, I have met with a reaction on brass which, so far as I know, has not before been recorded and of which this note is a preliminary notice. If a small piece of brass or a few brass turnings be covered with liquor ammonia, sp. gr. 0.880, in a closely fitted stoppered

bottle, and placed aside for a few days, it will be found that the ammonia has acted on the copper of the brass to such an extent as to produce a solution of a more or less characteristic violet colour, due to the presence of oxide of copper held in solution by ammonia. If this solution be still allowed to remain undisturbed for a few days longer free from contact with the air, this violet colour will gradually disappear, leaving a colourless solution, which, however, is no sooner brought in contact with the air by removing the stopper than the violet colour is reproduced, and by again stopping the bottle and leaving it aside the same reaction occurs and may be reproduced over and over again. The production of the violet colour from a colourless solution on exposure to the air does not seem to be the result of oxidation, as on opening the bottle in an atmosphere of carbonic acid the same reaction takes place.

I am at present working up this subject in the hope of finding in what state this colourless solution of copper exists.

Cullochfaulda Chemical Works,
Glasgow, November 10, 1879.

PROCEEDINGS OF SOCIETIES

CHEMICAL SOCIETY.

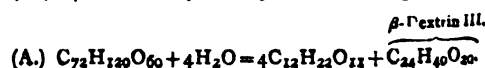
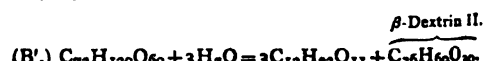
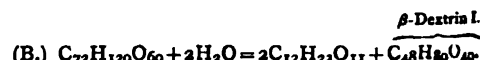
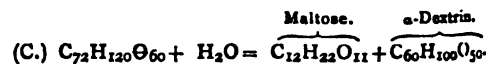
Thursday, November 6, 1879.

Mr. WARREN DE LA RUE, President, in the Chair.

THE minutes of the last meeting were read and confirmed. A list of presents to the library was read, and the thanks of the Society voted to the respective donors. The following certificates were read for the first time:—G. S. Allbright, W. J. F. Churchouse, M. Cochrane, W. R. Dunstan, J. J. Hummel, E. Hughes, T. S. Humpidge, F. Hatton, A. Leibius, R. Jones, H. F. Morley, H. Newton, J. Parette, G. Stallard, J. M. Wilson.

The PRESIDENT announced that the Council had expended about £300 in purchasing books for reference and circulation. He invited Fellows to offer suggestions as to the purchase of new books and the general improvement of the circulating library.

The SECRETARY then read a paper by C. O'SULLIVAN, "On the Transformation-products of Starch." This paper was originally presented to the President of the Société Chimique de Paris on June 18; extracts therefrom were read on July 4; but as the bye-laws of the Council exclude papers longer than eight printed pages, the author thought it desirable to bring the communication before the Society, especially as it appeared from papers published by MM. Musculus and Gruber that these chemists were, to a great extent, unacquainted with the previous work of the author. The author commences by reasserting that the molecule of starch under the influence of malt extract splits up in one of four ways:—



Other proportions of maltose and dextrin have been observed, but these are due to either the splitting-up of

the starch partially according to one equation, and partially according to one or more others, or to the further action of the active agents of malt extract on the dextrin first produced. The proportions of maltose and dextrin represented by the equation of MM. Musculus and Gruber belongs to the latter class. These chemists confirm the author's work as far as it relates to maltose, but have made a serious error (about 2 per cent) in calculating the specific rotatory power, as they have taken 220 m. m. of a 1 per cent solution of maltose to give a deviation of 13.5 divisions on the scale of the Soleil-Duboscq instrument. This number should be 13.75. MM. Musculus and Gruber point out at least three dextrins with different optical activities and cupric oxide reducing powers. The author declares the existence of four distinct dextrins to be highly probable, but states that all, when pure, have the same optical activity, but that none are reducing bodies. Soluble starch has the same optical activity as the dextrins, but like them does not reduce when pure. The author then gives details of the preparation and properties of soluble starch and the various dextrins, criticising the statements and methods of MM. Musculus and Gruber, as well as the various products obtained by them. In the second part the author investigates the action of malt extract on the transformed products. At first sight it would seem probable that the starch molecule breaks down first into maltose and α -dextrin; the latter is then converted into maltose and β -dextrin I., which in its turn forms maltose and β -dextrin II., &c. The author has made many experiments to elucidate this question, and finds that the above theory—viz., the breaking down of the starch molecule into dextrin molecules which become smaller at each step—is not supported by all the facts. Continual work with these bodies has forced the author to the conclusion that they are not a series of polymers, but rather a series of bodies of the same molecular weight, in which the difference of behaviour must be accounted for by a difference of relation in the arrangement of the molecules to one another, probably in solution alone. We may take it, continues the author, that the molecule of soluble starch, or the dextrin giving a purple with iodine, is simply $C_{12}H_{20}O_{10}$; but that these molecules in solution are arranged in groups of six sixes, all the groups being in an intimate state of tension one with another, so that the motion affecting one under certain conditions affects all under the same condition. In conclusion, the author states that the theory of splitting up and breaking down of the starch molecule, as represented by the equations, does not hold all the facts eliminated, and is not in accord with them. The theory, on the other hand, of the arrangement of the molecules in groups, all dependent one on the other, and capable, therefore, of undergoing a simultaneous movement, and the re-arrangement of these groups attendant upon the hydration of a definite proportion of the molecules in each of them, holds all the facts at present known, and is in perfect accord with all of them. The author is still engaged on the chemistry of the subject, but points out that the physics ought now to be studied, and the heat absorbed or eliminated during the different transformations determined; thus some idea of the character of the apparent work done will be gained.

Dr. ARMSTRONG then read a "Note on the Formula of the Carbohydrates." The simplest carbohydrates—dextroglucose and its isomerides—are closely related to mannite and dulcite, as shown by their conversion into one or other of these alcohols by the action of nascent hydrogen, and therefore they are derivatives of the paraffin, normal hexane. Of the three possible formulæ which may be assigned to a body formed from mannite or dulcite by the absorption of 2 atoms of hydrogen, that which represents glucose as being both an aldehyd and a penthydric alcohol,



appears to be the most probable, as it is the only one which accounts for the formation of saccharic or mucic acids from it on oxidation. The carbo-hydrates of

the cane-sugar group are probably related to the glucoses in the same way that ordinary ether is related to ethyl alcohol; moreover, it appears most probable that the simplest carbohydrate of the empirical formula $C_6H_{10}O_5$, i.e., dextrin, and if several exist, the lowest of them, bears a similar relation to the carbohydrates of the cane sugar group, and therefore has the formula $C_{24}H_{40}O_{20}$. Supposing starch to be a body of highly complex formula, the author is inclined to prefer this hypothesis to that advanced by O'Sullivan; there is very little doubt that it bears an altogether different relation to the lower carbohydrates; most probably the relation is of the character of that which obtains between aldehyd and its polymerides, i.e., the groups composing the starch molecule are partly held together by the COH groups, these groups being rendered capable of thus acting by their conversion into HC—O— groups. On this hypothesis a number of $C_6H_{10}O_5$ polymerides of varying molecular weight may be conceived to be capable of existing, and bodies such as inulin, glycogen, &c., are not improbable by intermediate terms in such a series. The fact that the dextrins obtained by the decomposition of starch under various conditions are very similar in their properties is not so difficult to reconcile with this hypothesis, if it be assumed, as appears probable, that the polymerisation and formation of higher terms in the series are attended with the expenditure of only a small amount of energy; in this case the polymerides would not, probably, differ greatly from each other in properties. It may be argued that as cane sugar and the dextrins are not cupric oxide reducing bodies, it cannot be assumed that they contain the group COH in their formulæ; probably, however, the power of reducing cupric salts is in no way connected with the presence of COH groups. In conclusion, the author considers that there is little doubt that our present theory of isomerism is insufficient and incapable of explaining the isomerism of mannite with dulcite and of the glucoses with one another. The Le Bel hypothesis appears at first sight to be applicable, but there is much evidence tending to show that this hypothesis does not furnish the entire solution of the problem.

The next paper was read by the SECRETARY, "On a New Method of Determining Sulphur in Coal," by TEIKICHI NAKAMURA, of the Engineering College, Tôkiô. The author criticises the methods generally used, the nitric acid and potassium chlorate method, the potassium hydrate method, and the method of fusion with alkali carbonates and nitre, and he finds them all more or less inaccurate or inconvenient. He recommends the following procedure:—3 or 4 parts of the mixed alkali carbonates, or of sodium carbonate, are intimately mixed with one part of coal in very fine powder in a large platinum dish. The mixture is heated at first very gently, a spirit lamp being used to prevent possible absorption of sulphur, instead of a Bunsen; the heat is then raised slowly without attaining that of visible redness until the surface becomes only faintly grey. No smoke or odorous gases should escape during the whole of the oxidation. The temperature is now raised to a faintly red heat for sixty minutes, at the end of which time the mass is perfectly white or reddish if iron be present. The mass is not to be stirred during the ignition. The residue is treated with water, filtered, and the sulphates determined in the ordinary way. The author quotes some results obtained by his process. To the paper is appended a note by Dr. Divers, stating that the above process was worked out some time before the CHEMICAL NEWS of January 17th was received, which contains a somewhat similar process by Mr. Pattinson, who estimates the sulphur by heating a mixture of coal and calcium hydrate in a muffle.

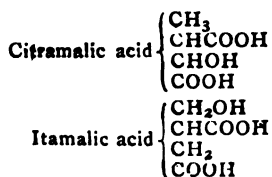
At the conclusion of this paper, Dr. GILBERT took the chair.

The next paper was read by the SECRETARY, "On the Bromine Derivatives of β -Naphthol," by A. J. SMITH.

The author prepared β -mono-brom-naphthol by adding to a tolerably concentrated solution of β -naphthol in glacial acetic acid, the theoretical amount of bromine mixed with an equal volume of glacial acetic acid; the bromine is added drop by drop, and the mixture kept cool. On standing colourless needles with an adamantine lustre crystallise out. These were purified and analysed. β -mono-brom-naphthol is soluble in alcohol, ether, and benzene, melts at 84° C., begins to decompose at 130° ; when oxidised by alkaline permanganate it yields orthophthalic acid. The author prepared β -tetra-brom-naphthol in a similar way: on oxidation it yielded monobromophthalic acid. In conclusion, the author discusses the bearing of his results on the constitution of these bodies.

The next paper was read by the SECRETARY, "Notes on the Dissociation of Ammonia Iron Alum," by J. S. THOMSON. When a dilute neutral solution of a ferric salt is heated, it dissociates, and a basic salt is precipitated. This basic salt is readily soluble in dilute sulphuric acid, if an acid of known strength be employed, and the quantity necessary to bring about the desired solution be known, an insight into the nature of the decomposition can be arrived at. The author refers to the previous work of Krecke, Wiedemann, Tichborne, Naumann, and others. Several series of experiments were made. Taking the mean of these, the quantity of sulphuric acid required to prevent dissociation for each addition of 10 c.c. of water, after 50 c.c., when ammonia iron alum equal to 0.1 gm. Fe_2O_3 is used, was found to be 0.0186 gm.; if 0.15 gm. Fe_2O_3 be used, the quantity of H_2SO_4 was 0.02479 gm. The addition of sulphate of ammonia equal to 0.05 ammonia necessitates an additional quantity of sulphuric acid equal to 0.05005 gm. Ammonia salts exert their influence when present as double salts. Ammonia alumina alum, the double sulphates of ammonia and magnesia, and of ammonia and zinc were used. Potash salts exert a still more powerful influence in promoting dissociation. A solution of ammonia iron alum containing more than 1 gm. in 14.37 c.c. does not dissociate on boiling. The dissociation begins in more dilute solutions, and increases regularly, as the above results prove, with each addition of water.

The next paper was read by the SECRETARY, "On α -Methyl Oxysuccinic Acid, the Product of the Action of Anhydrous Hydrocyanic Acid upon Aceto-acetic Ether," by G. H. MORRIS, Demargay (*Comptes Rendus*, lxxxii., 1337) has described an acid, obtained as above, under the name of oxypyrotartaric acid, as an unstable uncrystallisable syrup whose barium salt is decomposed by boiling with excess of water. The author, at the suggestion of Wislicenus, has prepared the body according to the directions of Demargay. The thick brown syrup thus obtained was purified by solution in water precipitated with lead acetate, the lead salt treated with sulphuretted hydrogen, the solution filtered, evaporated, treated with ether, until finally the ethereal solution, after standing a few days in the air-pump vacuum, deposited the pure acid in star-like groups of needles, deliquescent, and melting at 108° . The following salts were prepared and analysed:—The barium salt, which, when pure, did not decompose on boiling with water; the calcium, potassium, silver, lead, and copper salts, most of which are deliquescent. The author has investigated the action of fuming hydriodic acid and the products of the dry distillation of the acid. In conclusion, he discusses the constitution of some of the isomers of α -methyl oxysuccinic acid, and assigns the following formulae:—



and



The next paper was read by the SECRETARY, "On the Action of Phosgene on Ammonia," by H. J. H. PENTON. When these gases are mixed, a white neutral amorphous substance is produced, which has been shown by Regnault and others to consist of ammonium chloride, a substance identical or isomorphous with urea, and in addition small quantities of guanidine, cyanuric acid, &c. (Bouchardat). The author prepared some quantity of this white substance, and obtained a small quantity of guanidine, which he identified by its crystalline form, and some urea, which was identical with ordinary urea in its behaviour with hypochlorites and hypobromites (*Yours. Chem. Soc.*, July, 1878), and in other respects. The author concludes that either carbamide and urea are identical, or that symmetrical carbamids has not been obtained by this method.

The next paper was read by the SECRETARY, "On the Rehydration of Dehydrated Metallic Oxides," by C. F. CROSS. The author has obtained various anhydrous basic metallic oxides by igniting the precipitated hydrates. These oxides he exposed at the ordinary temperature to an atmosphere saturated with aqueous vapour. Rehydration occurs up to a definite limit of a molecular character, attended in most cases with a change of volume. He has investigated thus the oxides of aluminium, chromium, cobalt, iron, and copper.

The following paper was taken as read: "On Alizarin Blue," by G. AUERBACH. About eighteen months since a blue colouring matter was brought into the market as a substitute for indigo. It is now disused on account of its high price and its unstable nature when exposed to sunlight. The researches contained in this paper were finished in May, 1878. The author gives a *resumé* of previous work on the subject, and recommends the following method of preparation:—1 part of dry mono-nitro-alizarin, 5 parts of concentrated sulphuric acid, and 1 part of glycerin (sp. gr., 1.262), are mixed and heated gently. Reaction commences at 107° C., becomes violent, the temperature rising to 200° . Much frothing takes place, with evolution of sulphurous acid and acrolein. The whole mass, when frothing has subsided, is poured into water, boiled up and filtered, the residue being boiled out three or four times with dilute sulphuric acid. The mixed filtrates are allowed to cool, and the blue separates in brown crystals. These are purified by mixing with water and adding borax till the solution becomes brownish violet, the blue with the boric acid forming an insoluble compound. This residue is washed, decomposed with an acid, and the pure blue obtained as a violet silky paste. If required perfectly pure, it must be crystallised successively from its various solvents, high-boiling naphtha, amyl alcohol, and glacial acetic acid. When pure it forms brown shining needles, melting $268-270^{\circ}$. It has the formula $\text{C}_{17}\text{H}_{11}\text{NO}_4$. Salts were prepared and analysed, but the results were not satisfactory, as it was difficult to obtain them quite pure. Bromine derivatives were also prepared and examined. The action of chlorine, zinc dust, acetic anhydride, &c., have also been studied. The author discusses the constitution of the blue, and thinks it must be closely related to the aldehydines discovered by Ladenburg, which are formed when aromatic orthodiamides act upon aldehyds.

The Society then adjourned to November 20th, when the following papers will be read:—"A Chemical Study of Vegetable Albinism," Part II.; and "The Respiration and Transformation of Albino Foliage," by A. H. Church; "On the Estimation of Manganese Oxides," by S. Pierling; "A Contribution to the History of Putrefaction," by C. T. Kingzett; "Notes on Manganese Dioxide," by G. R. A. Wright and A. C. Menke.

PHYSICAL SOCIETY.

Ordinary Meeting, November 8, 1879.

Prof. W. G. ADAMS in the Chair.

THE meetings of this Society were resumed for the season on Saturday, November 8.

The first paper read was "On an Analogy between the Conductivity for Heat and the Induction Balance Effect of Copper-tin Alloys," by W. CHANDLER ROBERTS, F.R.S. Mr. Roberts traced a remarkable resemblance between a curve representing the induction balance effect of the copper-tin alloys published by him in June last, and the curve of Calvert and Johnston for the conductivity of heat, and, on the other hand, he showed that the induction curve does not agree with Matthiessen's curve for the electric conductivity of the same alloys. The author showed that the two alloys which occupy critical points of the curve, SnCu_3 and SnCu_4 , are of much interest. Possibly both are chemical combinations, and the wide difference in the position they occupy probably marks a difference of allotropic state. For the solution of such questions, however, Mr. Roberts considered that we might look with confidence to Prof. Hughes's beautiful instrument, which, he hopes, will also enable us to determine whether the relation between conductivity for heat and electricity is really as exact as it has hitherto been supposed to be.

As supplementary to this subject Dr. O. J. LODGE stated that he had compared the conductivity of six bars of the tin copper alloys, measured by the balance and by the Wheatstone bridge, and found them to agree very closely on the whole. The bridge results confirmed the resemblance traced by Mr. Roberts still more than the induction balance results.

Prof. HUGHES expressed his opinion that existing tables of metal-conductivity were erroneous. They disagreed among themselves, and the induction balance showed that it was impossible to get two pieces of the same metal exactly alike; hence the variation of specific conductivity results.

Prof. AYRTON stated that at a former meeting he had suggested that the electric inertia of the different specimens of metal tested might cause the difference between the results obtained by the Wheatstone bridge and the induction balance. Mathematical calculation had since led him to the conclusion that the inductive effect is not proportional to the resistance of the metal tested, but to a power or exponential of the resistance.

Prof. HUGHES replied that as the inductive effect of the metal was destroyed by cutting it so as to interrupt the circuit in it, it was reasonable to suppose that the said effect was due to induced currents circulating in the metal, and therefore was proportional to the conductivity of the metal.

Capt. ARMSTRONG exhibited a standard Daniell cell formed of porcelain vessel with a porous partition dividing it into two compartments. In one the zinc plate was immersed in a solution of sulphate of zinc; in the other the copper plate in a solution of sulphate of copper. To use the cell as a standard, it was only necessary to connect the two liquids by a cotton string moistened with water. This arrangement prevented mixing of the liquids, as the string could be withdrawn after use. The resistance was high, but it was a constant standard of electromotive force.

Prof. GUTHRIE mentioned that Prof. Pirani, of Melbourne, in a letter to him had signalled the fact that when a dilute acid was being electrolysed, the positive electrode, if made of iron, became incandescent below the surface of the liquid. Prof. Guthrie had found this to be true not only for iron, but for other metals, and that it could hardly be due to oxidation, because it took place not only at the cathode or positive electrode, where oxygen was evolved, but also at the anode, where hydrogen was evolved. The

incandescence appeared to him to be due rather to resistance. The author exhibited his experimental results, which he did not doubt had already been obtained by Prof. Pirani himself. The positive electrode when immersed in the electrolyte was seen to get red-hot and to wobble about. As the liquid heated the red glow became fainter. The negative electrode, on the other hand, emitted a bright light, accompanied by a sputtering noise. The light was tinged with the characteristic colour of the flame of the metal of which it was composed; in the case of a copper electrode, for example, it was greenish. These effects were shown by Prof. Guthrie with iron, copper, and platinum electrodes, in dilute sulphuric and dilute nitric acid.

In reply to Prof. Adams, Prof. GUTHRIE said he had not yet examined the flame by the spectroscope; and in reply to Prof. Foster, he stated that the battery power used was 50 Grove's cells. He asked for suggestions as to the true cause of the phenomenon.

CORRESPONDENCE.

NORMAL SOLUTIONS.

To the Editor of the Chemical News.

SIR,—What is a normal solution? Unfortunately the term seems to be used in two distinctly different senses. By Sutton, whose book is best known to English chemists, the term is employed to signify a standard solution containing one equivalent of the active constituent in 1000 measures of water. I have not Sutton's work before me, but he uses the term in the same sense as I have done in my "Commercial Organic Analysis," in which a normal solution is defined (page 20) as "one containing in 1000 cubic centimetres such an amount of the active constituent as will combine with, replace, or oxidise 1 grm. of hydrogen." According to this definition, the following are the strengths of typical normal solutions:—

	Grms. per litre.
Normal caustic soda contains..	$\frac{\text{NaHO}}{40} = 40$
" carbonate sodium ..	$\frac{\text{Na}_2\text{CO}_3}{2} = 53$
" baryta water ..	$\frac{\text{BaO}}{2} = 76.5$
" hydrochloric acid ..	$\frac{\text{HCl}}{36.5} = 36.5$
" sulphuric acid ..	$\frac{\text{H}_2\text{SO}_4}{2} = 49$

Similarly the following solutions are decinormal:—

Decinormal silver nitrate contains ..	$\frac{\text{AgNO}_3}{10} = 17.0$
" mercuric chloride ..	$\frac{\text{HgCl}_2}{20} = 13.55$
" potassium permanganate ..	$\frac{\text{KMnO}_4}{50} = 3.162$

The advantage of the above arrangement is that all "normal" solutions are of exactly corresponding strength, and those similar in nature may be substituted one for another.

By Fleischer, a normal solution is understood to mean a liquid containing the atomic weight in grammes in 1000 c.c. As he employs the old atomic weights, in most cases the above strengths are called normal; but in the translation of his book by Mr. M. M. Pattison Muir, the doubled atomic weights are used with the consequence of very serious obscurity. Thus, while normal caustic potash would be held to contain 39.1 grms. of K per litre, we are told that "a normal solution of potassium carbonate is prepared by dissolving 138.2 grms. of potassium carbonate in 1000 c.c. of distilled water" (p. 37). Hence such a solution contains 78.2 of K, and is twice as strong

as the caustic alkali, or as the normal potassium carbonate of the original German.

The confusion is still greater when permanganate comes to be used. In the original German, Fleischer calls a solution of 32 grms. of potassium permanganate in 1 litre of water, $\frac{1}{2}$ normal. It is true that this solution is absurdly strong, and is capable of doing ten times the work attributed to it by the author. Mr. Muir has corrected this in an erratum, but in the text of his translation he actually doubles the strength of the above solution, recommending one containing 64 grms. per litre. Hence we have the following confusion in nomenclature.

According to Fleischer, $\frac{1}{2}$ normal permanganate contains 32 grms. per litre, or, if this large amount be due to printer's error, 3.2 grms. per litre.

According to his translator, Muir, $\frac{1}{2}$ normal permanganate contains 64 grms. per litre, though it is doubtful, owing to the erratum, whether he would not consider that the solution so described ought to contain 64 grms. per litre.

According to Sutton, $\frac{1}{2}$ normal permanganate contains 6.324 grms. per litre.

That the nomenclature of a standard solution should vary according as the writer employs the old or doubled atomic weights must be held to be highly inconvenient, to say the least of it, though Sutton (compare fit with later editions) has wholly avoided any such source of confusion. According to the principle adopted by Mr. Muir, even the employment of the same atomic weights will not always save us, for a standard solution of permanganate will be differently called according as its employer considers the salt in solution to be KMnO_4 , or disregards its isomorphism with K_2CO_3 , and prefers to write it $\text{K}_2\text{Mn}_2\text{O}_8$.

One other instance of this confusion of terms. Mr. J. B. Hannay, in his paper on the "Volumetric Estimation of Cyanides," calls a solution containing 27.1 grms. of mercuric chloride to the litre "*decinormal*," although a litre would correspond to *two-tenths* of an atom of cyanogen.

As may be gathered from the foregoing remarks, I myself prefer Sutton's interpretation of the term "normal" as applied to standard solutions, but I write this letter not so much to advocate my own views, as to point out the confusion which exists, in the hope that chemists may avoid using a well known term in a different sense from that in which it has been commonly employed.—I am, &c.,

ALFRED H. ALLEN.

Sheffield, Nov. 10, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances de l'Académie des Sciences. No. 16, October 20, 1879.

Observations of Magnetic Declension, Inclination, and Horizontal Intensity in the Basin of the Mediterranean.—M. de Bernardière.—Tables of results.

The Saccharimeter of Laurent.—L. Laurent.—A description of this instrument with figures. It is said to yield more light and more distinctness, and the reflections in the tubes are suppressed.

No. 17, October 27, 1879.

Galvanic Oxidation of Gold.—M. Berthelot.—Grotthus observed the dissolution of a gold wire used as a positive pole in sulphuric acid traversed by the current. M. Chevreul having suggested that this effect might be

due to the formation of persulphuric acid, certain experiments were undertaken to solve the question. The statement of Grotthus was confirmed. Gold was rapidly dissolved, even in dilute sulphuric acid, a part being reprecipitated at the negative pole. Nitric acid also attacks gold under the same conditions, but phosphoric acid and potassa have no appreciable action. The effect of sulphuric and nitric acids upon gold is not due to ozone, since oxygen charged with ozone has no action upon gold in presence of water, whether pure or mixed with sulphuric or nitric acid. Persulphuric acid, even when containing oxygenated water, does not attack gold.

Decomposition of Hydro-selenic Acid by Mercury.—M. Berthelot.—Hydro-selenic acid gas preserved in bottles for some years, at common temperatures and in contact with mercury, is slowly decomposed, forming mercury selenide.

Critical Reflections on the Experiments Concerning Human Heat.—M. Hirn.

Specific Heats and Melting-points of Various Refractory Metals.—J. Violle.—The specific heat of iridium, like that of platinum, increases regularly with the temperature. The mean specific heat of gold varies little up to 600°, and then increases sensibly on approaching the melting-point. The melting-points given are:—Silver, 954°; gold, 1035°; copper, 1054°; palladium, 1500°; plat num, 1775°; iridium, 1950°.

Chloride of Lime Battery.—A. Naudet.—The battery has for its positive electrode a plate of zinc, and for its negative electrode a plate of coke surrounded with fragments of coke. The zinc is placed in a solution of common salt; the coke is surrounded with chloride of lime, in a vessel of biscuit ware or of parchment paper.

Combinations of Hydrogen Phosphide with the Hydracids, and their Heats of Formation.—J. Oger.—The author has studied the hydrochlorate, hydrobromate, and hydriodate of hydrogen phosphide. The formation-heat of the latter compound is smaller than that of ammoniacal gas, which is conformable with analogies. The formation-heats of the ammoniacal salts are also notably greater than those of the phosphoretted compounds.

On Erbium.—P. T. Clève.—See CHEMICAL NEWS, vol. xl, p. 224.

Commercial Trimethyl-amin.—E. Du villier and A. Buisine.—The authors re-affirm their former conclusions concerning the trimethyl-amin prepared by the process of M. Vincent, in which they have further detected ethyl-amin and oxamids.

Ordinary Cellulose.—M. Franchimont.—The author adds to a mixture of 1 part of cellulose and 4 parts of acetic anhydride a little sulphuric acid, when a brisk reaction is set up and the cellulose disappears, whilst the liquid becomes coloured. The whole is then thrown into a large excess of cold water, which gives a copious white precipitate. The precipitate is washed in cold water and dried in the air, and is next introduced into alcohol, which dissolves a part and turns slightly yellow. It is filtered, washed in alcohol, and the residue dissolved in boiling alcohol. The solution deposits acicular crystals, composed of $\text{C}_{46}\text{H}_{54}\text{O}_{27}$. It seems to be an eleven times acetylated derivative of a triglucose, $\text{C}_{18}\text{H}_{32}\text{O}_{16}$.

On Glucose.—M. Franchimont.—The author describes an octacetytic diglucose, which has entirely lost the tendency to oxidation present in glucose.

Researches on Colour-blindness.—J. Macé and W. Nicaï.—The authors have sought to obtain comparative measurements between the quantities of light perceived in the different parts of the spectrum by the colour-blind on the one hand, and by the normal eye on the other. A red glass which scarcely lessens the vision of the normal eye diminishes remarkably that of a red colour-blind eye.

Justus Liebig's Annalen der Chemie,
Band 199, Heft 1.

Separation of the Heavy Metals of the Ammonium Sulphide Group.—C. Zimmermann.—Inserted at length.

Coto-barks and their Characteristic Constituents.—J. Jobst and O. Hesse.—This bark is probably derived not, like the true cinchonas, from a Rubiaceae plant, but from one of the Laurineæ or the Terebinthinaceæ. It contains a principle, which the authors name cotoin, $C_{22}H_{18}O_6$. Its reactions and derivatives are described at some length.

The Non-existence of Pentathionic Acid.—W. Spring.—The substance hitherto known as pentathionic acid is simply tetrathionic acid. On passing sulphurous acid and sulphuretted hydrogen into water, the former being maintained in slight excess, a liquid is obtained which distinctly decolorises indigo, and which is consequently the hydrosulphurous acid of Schützenberger.

Certain New Basic Salts of Mercury Sulphide.—W. Spring.—The author has analysed the compound produced on adding tetrathionic acid to an aqueous solution of mercurous nitrate—a yellow amorphous flocculent body, insoluble in water. It is a trithio-basic mercury sulphate. When treated with nitric acid it is gradually converted into a white body—mono-thio-basic trisulphate.

Bromoxyl Derivatives of Benzol.—R. Benedict.—The author describes the formation of tribrom-phenol-brom, its behaviour with alcohol, with tin and hydrochloric acid, with aniline and phenol, and with sulphuric acid.

Normal Paraffins.—C. Schorlemmer.—The results of this investigation show that during the action of chlorine upon normal hexan from mannite, a secondary chloride is formed in larger quantity along with the primary. The normal paraffins of mineral oil have always a higher specific gravity than those from other sources.

Suberic and Azelaic Acids.—R. S. Dale and C. Schorlemmer.—On treating cork with 4 parts of nitric acid at 1.30, the clear liquid contained in addition to suberic acid abundance of azelaic acid, oxalic acid, and other acids not yet further examined, and smeary nitrogenous compounds. Suberon on treatment with nitric acid yields an aldehyd of α -pimelic acid. The pimelates of silver, calcium, and barium are described.

Glucoside of White Mustard Seed.—H. Will and A. Laubenheimer.—Sinabin has the composition— $C_{30}H_{44}N_2S_2O_{16}$.

The authors examine the behaviour of this compound with silver nitrate, mercuric chloride, and myrosin.

Chemiker Zeitung.
No. 42, 1879.

At the Congress of Naturalists and Physicians Prof. Engler discoursed on the dangers of petroleum, and exhibited his new instrument for ascertaining its safety. An induction spark is used to ignite the vapours given off on heating the oil to 35°. Prof. Schröder spoke on the stere of chlorine and oxygen, and Von Babo on determining the density of very small quantities of gas. Dr. Dabner (Döbner?) discoursed on the action of benzoyl-chloride and benzo-trichloride upon the phenols. Dr. Engler read a paper on the action of metals upon petroleum, and Prof. Volhard spoke on the compounds of manganese. For its detection he heats nitric acid and red-lead to ebullition, and adds drop by drop the dilute solution under examination. If manganese is present there is formation of potassium permanganate. Prof. Böttger spoke on the explosions sometimes produced by the action of sodium on water. He considers that the sodium peroxide formed gives off oxygen, which combines with the liberated hydrogen. Dr. Witt described the constitution of the gases evolved by the action of nitric acid upon arsenious

acid. Nölting, of Geneva, spoke on the formula of naphthalene, and Dr. Hesse closed the sectional meeting with a paper on quinhydrone. On the 23rd September, Prof. Jäger, of Stuttgart, gave, in the Conference Hall of the Gymnasium, a demonstration of his method of examining odours (neural-analysis) and on the odours themselves. At the last general meeting the professor spoke on the influence of the temper. Amidst boisterous merriment he appealed to all mothers present if children, when in a good humour, did not give off a pleasant odour, but a disagreeable one when cross or fretful. The lecturer ended amidst hisses. The number of members of the Congress this year was 1700.

Action of Bromine upon Cane-Sugar.—O. Griesshammer.—The products are a gluconic acid, a carbohydric resembling fructose, and a gum-like body. The acid is merely an isomer of the gluconic acid of Hlasiwetz.

Volumetric Determination of Magnesium.—Dr. H. Precht.—The author precipitates the soluble magnesian salts with potassa lye of known strength, and determines the excess of the latter with standard acid.

Apparatus and Methods for Testing Petroleum.—Dr. Skalweit.—The author gives the preference to the petroleum pyrometer of Sintenis.—*Hann. Monatschrift*.

Simple Method of Testing Petroleum.—Dr. L. Janke and Dr. A. Barth.—The authors heat in the water-bath, raising the temperature to 100° F. in ten minutes. At 95° they stir the oil, and at 100° F. they bring the flame of an ordinary match in direct contact with the liquid.—*Hann. Monatschrift*.

MISCELLANEOUS.

The Royal Institution.—The Christmas Lectures, adapted to a juvenile auditory, will this year be given by Professor Tyndall, the subject being "Water and Air."

Foundation of a New Chemical Chair.—Mr. Mark Firth has signified his wish to found a Chair of Chemistry in connection with Firth College, Sheffield. He proposes to invest a sum sufficient to produce an annual income of £150, and this, together with the fees of students, will amount, it is believed, to a handsome sum. Mr. Firth proposes that the appointment shall be first filled by Dr. Carnelley, of Owens College, Manchester. Mr. Firth built the College bearing his name, contributing at the same time £20,000 towards the general endowment fund.—*The Medical Press and Circular*.

NOTES AND QUERIES.

Polarimeter.—I should be obliged if you would be good enough to inform me through the *CHEMICAL NEWS* the best form of polarimeter for determining the amount of sugar in saccharine fluids, and the rotary power of other liquids.—J. C. H.

MEETINGS FOR THE WEEK.

WEDNESDAY, Nov. 19th.—Society of Arts, 8. First Ordinary Meeting. Opening Address of the 126th Session, by Lord Alfred S. Churchill.

Society of Public Analysis, 8. "On the Mineral Constituents of Cinnamon and Cassia," J. Henner. "On the Determination of Carbonic Acids in Carbonates," G. W. Wigner. "On the Examination of Coffee," A. H. Allen. "Note on Arsenic in Playing-cards," Dr. Waller. "Note on the Discrimination of Starches by means of Polarized Light," Dr. Tripe.

THURSDAY, 20th.—Chemical, 8. "A Chemical Study of Vegetable Albumin: Part II., Respiration and Transpiration of Amino Acids," A. H. Church. "Estimation of Manganic Oxide and Potassium Bichromate," Spencer Pickering. "Contributions to the History of Putrefaction," C. T. Kingzett. "Notes on Manganese Dioxide," C. A. Alder Wright and A. E. Menke. Royal, 8.30.

SATURDAY, 22nd.—Physical, 3. "On a Retention Image Photometer," Dr. F. Guthrie.

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INSTITUTE OF CHEMISTRY.

The President has offered Two Prizes of £50 each for the two best original investigations involving Gas Analysis. These Prizes will be open to Associates, and to all persons (except Fellows of the Institute) who shall before the 31st December next have qualified for the Associateship in all respects short of passing the prescribed practical examination, and successful competition for these prizes will be accepted in lieu of such practical examination.—Further information may be obtained on application to the Secretary, Mr. C. E. GROVES, Somerset House Terrace, W.C.

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THE PERIODIC LAW OF THE CHEMICAL ELEMENTS.

By D. MENDELEEF.

(Continued from page 232.)

EVEN as, up to the time of Laurent and Gerhardt, the words "molecule," "atom," and "equivalent" were used one-for the other indiscriminately in the same manner, so now the terms "simple body" and "element" are often confounded one with the other. They have, however, each a distinct meaning, which it is necessary to point out, so as to prevent confusion of terms in philosophical chemistry.

A "simple body" is something material, metal or metalloid, endowed with physical properties, and capable of chemical reactions. The idea of a molecule corresponds with the expression of a "simple body;" a molecule is made up of one atom, as in the case of Hg, Cd, and others; or of several atoms, such as S₂, S₆, O₂, H₂, Cl₂, P₄, &c.

A simple body is able to show itself under isomeric and polymeric modifications, and it is only distinguished from a compound body by the homogeneity of its material parts. But in opposition to this, the name of "element" must be reserved for characterising the material particles which form simple and compound bodies, and which determine their behaviour from a chemical and physical point of view. The word "element" calls to mind the idea of an atom; carbon is an element; coal, diamond, and graphite are simple bodies.

The principal end of modern chemistry is to extend our knowledge of the relations between the composition, the reactions, and the qualities of simple and compound bodies, on the one hand; and, on the other hand, the intrinsic qualities of elements which are contained in them; so as to be able to deduce from the known character of an element all the properties of all its compounds.

For example, saying that carbon is a tetratomic element, is making known a fundamental property which appears in all its combinations.

The elements count among their properties, which can be measured exactly, their atomic weight, and the power of showing themselves under the form of different compounds.

Alone amongst their properties, the two above mentioned bring in their train a number of facts. The last has given rise to a special theory on the atomicity (valency) of elements. Amongst the other properties of elements which influence the character of bodies, the physical properties (such as cohesion, capacity for heat, coefficient of refrangibility, spectral phenomena, &c.) have been up to the present time too incompletely studied for us to be able to generalise them in a rigorously philosophical manner. What we know of these properties is still insufficient and defective in comparison with our knowledge on the atomic weights and the atomicity of elements. However, it has already been often noticed that the physical properties depend one on another, that the atomic weights, and principally the molecular weights of compounds, are equally in intimate relation with them. It is principally the fact, that it is easy to measure these properties exactly, that has induced us to make these comparisons. It is by studying them, more than by any other means, that we can conceive the idea of an atom and of a molecule. By this fact alone we are enabled to perceive the great influence that studies carried on in this direction can exercise on the progress of chemistry.

The above-mentioned measurable properties are by no means the only ones possessed by the elements. They have beyond these a series of properties which have not yet been able to be measured, but which still contribute to their recognition. These last have received the name of chemical properties. Certain elements do not combine with hydrogen; they have, according to the recognised term, a basic character, or, in other words, they absorb oxygen and form bases; they form salts when combining with chlorine; other elements (called acidifying elements) do combine with hydrogen; with oxygen they only form acids, and with chlorine only chloranhydrides. Thirdly, there are elements which form the link between the first and second classes; and fourthly, there are elements which in their forms of higher oxidation have an acid character, and when less oxidised a basic character. Science does not as yet possess any process by which these properties can be measured, but still they are counted among the number of qualitative characteristics which distinguish the elements. Further, these last-named elements possess properties which determine the greater or less stability of compounds; these, again, are chemical properties. It is in this manner that some elements can unite with all the others in compounds capable of being decomposed with a relative facility, while we cannot obtain analogous decompositions in the corresponding compounds of other elements. Not being susceptible of exact measurement, the above-mentioned chemical properties can hardly serve to generalise chemical knowledge; they alone cannot serve as a basis for theoretical considerations. However, these properties should not be altogether neglected, as they explain a great number of chemical phenomena. It is known that Berzelius and other chemists considered these properties as being among the principal characteristics of elements, and that it is on them that the electro-chemical system was based.

As a general rule when we study the properties of elements, bearing in mind practical conclusions and chemical previsions, it is necessary to give equal attention to the general properties of the other bodies of the group, and to the individual properties of the given element in that group; it is only after such comparative studies, and laying stress on an accurately measurable property, that we can generalise the properties of an element. The atomic weight furnishes us now, and will long continue to furnish us, with a property of this nature; for our conception of the atomic weight has acquired an indestructible solidity, above all latterly, since the use of Avogadro's and Ampère's law: thanks to the efforts of Laurent, Gerhardt, Regnault, Rose, and Cannizzaro, one can even state boldly that the notion of the atomic weight (considered as the smallest part of an element contained in a molecule of its compounds) will remain without change, whatever may be the modifications that the philosophical ideas of chemists may undergo. The expression atomic weight* implies, it is true, the hypothesis of the atomic structure of bodies; but, then, we are not here discussing denomination, but a conventional idea. It seems that the best method of extending our chemical knowledge would be to elaborate the correlations between the proportions of elements and their atomic weights. It is thus that we should obtain the most natural and most fruitful results in the extension of the study of elements. To determine this dependence seems to me to be one of the principal tasks of future chemists; for this problem has the same philosophical importance as the study of the conditions of isomerism has. In the present memoir I shall try and show the already mentioned relation between the atomic weights of elements and their other properties, particularly the faculty of giving different forms of combination.

This last faculty has already been carefully experimented on; a still more precise expression has recently

* By replacing the expression of atomic weight by that of elementary weight, I think we should, in the case of elements, avoid the conception of atoms.

been found for it, in the theory relative to the limits of chemical combinations, to the atomicity of elements, and to the manner of attachment of atoms in the molecules. It is known that Dalton called combinations in multiple proportions the mode of combination of an ideal element, R, with other elements (of the form RX , RX_2 , RX_3 , &c.); Gerhardt called them types; they are now used for fixing the atomicity of elements.

The incompleteness which exists in the theory now accepted, with regard to the atomicity of elements, arises from the fact that the opinions of chemists do not coincide in respect to elements such as Na, Cl, S, N, P, Ag; some consider the atomicity as an invariable property of atoms, while others affirm the contrary. The uncertainty in the ideas on atomicity come principally from the novelty of their introduction into Science, and from this—that they include the hypothesis of the union of elements by parts of their affinity. It also arises, according to my idea, from the fact that we only study the forms of combination, without comparing these forms with the other properties of the elements. The gaps which I have just pointed out in the theory of combinations—gaps produced by the doctrine actually accepted on the subject of the atomicity of elements—are at their widest, as I shall point out further on, if the study of the principal properties of elements is based on the atomic weights.

Since the year 1868, the year in which the first part of my work "Principles of Chemistry" appeared, in the Russian language, I have been endeavouring to solve this problem. In this paper I take the liberty of making known the results obtained up to the present time in my researches in that direction.* The formation of natural groups, such as the haloids, the metals of the alkalis and alkaline earths, the bodies analogous to sulphur, to nitrogen, &c., furnished me with the first opportunity of comparing the different properties of the elements with their atomic weights. In the beginning we only arranged in groups the elements which resemble one another in several respects, but later on several experimentalists—notably Gladstone, Cooke, Pettenkoffer, Kremers, Dumas, Lenzen, Odling, &c.—observed that the atomic weights of the different members of these groups had a simple and regular relation to each other. The discovery of these relations led to the comparison of the members of different groups with the homologous series, and, later on, to the conception, in a chemico-mechanical manner, of the complex nature of atoms, which has been held as reasonable by the greater number of chemists, but up to the present time it has not received any definite name.

All the relations observed between the atomic weights of elements have not yet led to any logical conclusion or chemical prevision, on account of the gaps in them. This may be the reason why they have not acquired the right of being generally recognised in Science.

First. Nobody that I know of has, up to the present, prepared any comparative table of the natural groups, and the observed relations between the different members of groups have remained without any connection or explanation. Concerning this subject, in 1859, Sirecker rightly said,† "It is hardly possible that the relations noticed between the atomic weights of elements which resemble each other in their chemical properties should be purely accidental. However, we must leave to future research the discovery of the 'regular' relations which are betrayed in these numbers."

Secondly. Only small variations in the magnitude of the atomic weights of some analogous elements (Mn, Fe, Co, and Ni,—Pd, Rh, Ru,—Pt, Os, Ir) have been observed. Therefore we were only authorised in saying that the analogy of elements was connected either by approximate agreement or by the increasing amount of their atomic weights.

* In relation to some historical and polemical observations on this question, see the *Berichte der Deutschen Chemischen Gesellschaft*, 1871, p. 248.

† "Theorien und Experimente zur Bestimmung der Atomgewichte der Elemente," p. 146.

Thirdly. Nobody has established any theory of mutual comparison between the atomic weights of unlike elements, although it is precisely in connection with these unlike elements that a regular dependence should be pointed out between the properties and the modifications of the atomic weights. The facts published up to now, being too isolated, could not cause any progress in the philosophical development of chemistry; however, they contain the germs of important additions to chemical science, especially as concerns the nature, to us mysterious, of elements.

In the term *periodic law* I designate the reciprocal relations between the properties and the atomic weights of elements. Later on I shall develop the relations which are applicable to all the elements: they are shown in the form of a periodic function.

(To be continued.)

ON THE VOLATILITY OF PLATINUM IN CHLORINE GAS AT HIGH TEMPERATURES.*

By SEELHEIM, of Utrecht.

SOME years ago the author made the curious observation that, when platinum foil is kept at a red heat in dry chlorine gas, the metal gradually volatilises, and in a colder part of tube gets re-deposited as a sublimate, consisting of measurable crystals of the regular system. The crystalline nature of the sublimate proves that the metal must have travelled through the tube as a vapour. In order, however, to make quite sure of this important result, the author, quite lately, repeated the experiment in a modified form, consisting in this, that he heated a quantity of platinum chloride in a porcelain flask to bright redness. The flask was allowed to cool and then cautiously broken up, when the metallic platinum due from the chloride was found, not at the bottom of the flask, but somewhere higher up at the sides, in the form of a crystalline sublimate. This, the author says, confirms what was observed some time ago by Troost and Hautefeuille, who found that platinum chloride, when heated to 1400° C. in a porcelain flask and allowed to cool, suddenly gives a deposit of $PtCl_2$, while when cooled down gradually it yields only Cl_2 and metallic platinum.

The volatilisation of the platinum must be owing to a chemical cause. We must assume that the metal, when heated in chlorine gas, passes, at least, temporarily, into a volatile chloride, which, at lower temperatures, breaks up again into chlorine gas and metallic platinum. If this chloride is $PtCl_2$, and $PtCl_2 = 1$ molecule, its vapour should occupy the same volume as the " Cl_2 " contained in it, and consequently Victor Meyer's experiments prove, as before, that " Cl_2 " at high temperatures dissociates into two or more molecules. But it is more natural to assume that the $PtCl_2$ vapour has only a transitory existence, being continually formed and re-decomposed into Pt vapour and chlorine. At any rate, Victor Meyer's results do not prove that chlorine gas (at 1500° or so) undergoes dissociation, because what he operated upon was not pure chlorine but chlorine contaminated with volatilised platinum.

So far Seelheim. In my own opinion the most natural interpretation of Seelheim's and Meyer's results is to assume that $2(Pt + Cl_2)$, produced by the decomposition at a dull red heat of $2PtCl_2$ at higher temperatures, associate into $2PtCl + Cl_2 = 6$ vols. (i.e., 6 times the volume of $\frac{1}{2}H_2$) = $\frac{1}{2}$ times the volume the $2Cl_2$ present at the lower temperature. Or else we may assume that the atomic weight of platinum is $pt = \frac{1}{2}Pt$, and that, what on starting was $2pt_2Cl_2$ (solid), at higher temperatures got transmitted thus:—

* A condensed translation of an article in the last number of the *Berichte der Deutschen Chemischen Gesellschaft*. Communicated by Prof. Dittmar.

	$2\text{ptCl}_2 = \text{pt}_4 + 2\text{Cl}_2 = 2\text{ptCl} + \text{pt}_2\text{Cl}_2$
Ord. temp.	500° 1500°
Vol. =	0 0+4 4+2
or—	0 4 6

Meyer's discovery, then, appears to be a mistake, but it is one of those mistakes that could only have been committed by a great experimenter like him.

W. D.

Anderson's College, Glasgow, Nov. 17, 1879.

ON THE NASCENT STATE OF BODIES.

By Dr. T. TOMMASI.

In a recent number of the *CHEMICAL NEWS* (vol. xl., p. 184) Dr. Phipson observes that it is not logical to found an entire system on a single fact. In this he is perfectly right, and I entirely agree with him. I would, however, remind him that my thermic theory of nascent hydrogen does not rest on a single fact but on many. For more than two years I have been engaged in ascertaining whether the reductive properties peculiar to hydrogen when it comes out of a combination are due to an allotropic state of hydrogen, such as the nascent state, or to ordinary hydrogen in different thermic conditions. I have studied most of the reductions effected by hydrogen, and generally attributed to a nascent state such as—

- Reduction of silver chloride, bromide, and iodide.
- " " chloric acid and chlorates.
- " " potassium perchlorate.
- " " ferric chloride.
- " " nitrates.
- " " chloral.

The following are the results I obtained:—

(1.) Silver chloride in suspension in dilute sulphuric acid was treated with sodium amalgam. The experiment, which lasted forty minutes, was made in the dark. The silver chloride remained perfectly white, and consequently had undergone *no reduction*, and the liquid separated from the silver chloride contained no trace of sodium chloride. It was the same with silver bromide and iodide. It follows, moreover, that nascent hydrogen ($\text{Zn} + \text{SO}_4\text{H}_2 + \text{Aq}$) does not reduce silver chloride unless the chloride is in contact with zinc. This reduction, then, can only be attributed to the action of the zinc, and not to that of hydrogen. On the other hand, an electric current will decompose silver chloride in suspension in dilute sulphuric acid. How are we to explain the fact that silver chloride which resists the action of nascent hydrogen resulting from the decomposition of water by sodium amalgam can be reduced by hydrogen equally nascent, but which results from the decomposition of water by the battery?

(2.) A solution of potassic chlorate was acidulated with sulphuric acid, and then divided into two parts, one of which was treated with zinc, the other with sodium amalgam. The reaction was stopped at the same time, when there was still some sodium amalgam and some zinc, and before the sulphuric acid was completely neutralised. The two solutions were filtered, and silver nitrate added. The solution which had been treated with zinc gave rise to a very abundant precipitate of silver chloride, whilst the other, which had been treated with sodium amalgam, remained perfectly limpid. Chloric acid, sodium, barium, copper, and mercury chlorates undergo no reduction by sodium amalgam when in neutral, alkaline, or acid solution. The hydrogen proceeding from the electrolysis of water does not easily deoxidise potassium chlorate.

(3.) M. Guetta, in a work on the reduction of nitrates, has arrived at conclusions precisely similar to those I have arrived at with regard to chlorates. I had foreseen most of the results obtained by M. Guetta, and was glad to see them confirmed by his experiments.

(4.) M. Papasogli has recently discovered that a solution of nickel, to which potash and potassium cyanide were

added, acquires, by the action of the zinc, a beautiful red tint, whilst at the same time hydrogen is set free. If magnesium takes the place of zinc in this experiment—or, better still, if the magnesium-platina couple is used—the evolution of hydrogen is always observed, but the colouration no longer takes place. On the other hand, the double cyanide of potassium and of electrolysed nickel is coloured red at the negative pole. In these cases the red compound is either produced by the metal or nascent hydrogen. In the former case the red tinge would not be produced by the action of hydrogen resulting from the decomposition of water by the battery, and in the second case it would manifest itself, because the hydrogen set free by the magnesium-platina couple is equally nascent as that developed by the zinc or by the electric current.

(5.) M. Sergius Kern (*CHEMICAL NEWS*, vol. xxxiii., p. 112) observes that magnesium is rapidly dissolved by ferric chloride, ferric hydrate being formed. This paper was afterwards reproduced in the *Bulletin de la Société Chimique de Paris*, vol. xxvi., p. 338, but the editor of that journal remarks that it does not appear probable that ferric hydrate is produced when hydrogen is evolved unless air is admitted. I repeated that experiment, excluding the air, and using water that had first been boiled; but the results that I obtained were exactly identical with those of M. Sergius Kern. It is the same when sodium amalgam is substituted for magnesium.

(6.) According to Mr. C. Stahlschmidt (*Pogg. Annalen*, vol. cxxviii., p. 416) it would seem that the nascent hydrogen which results from the decomposition of water by powdered zinc at the ordinary temperature changes potassium nitrate into nitrite, red cyanide into yellow cyanide, reduces indigo and the iodates, but, most remarkably, *does not reduce the nitrates*.

(7.) Prof. De Wilde, of Brussels, has observed that whilst sodium amalgam reduced potassium bromate, it had no action on potassium chlorate. How, then, without recurring to my theory, would Dr. Phipson explain the fact that potassium perchlorate, which resists the action of sixteen sources of nascent hydrogen, can be reduced by sodium bisulphite? And why does the nascent hydrogen proceeding from the action of zinc on dilute sulphuric acid reduce the potassium chlorate while it has no action on the perchlorate? In the last two cases is it not always in the nascent state that the hydrogen is set free? and is not the medium in which it is produced the same? If, then, the properties of nascent hydrogen were inherent to that special state of the gas the same reactions ought always to be obtained; but the few experiments I have just quoted, and many others which I pass over in silence, prove, on the contrary, that the reductive power of nascent hydrogen varies according to the chemical reaction which produces it. And if this gas in the nascent state possesses greater affinity than in the natural state, it is solely due to the fact that the hydrogen the moment it issues from a combination is found to be accompanied by the whole quantity of heat produced during the setting free of the hydrogen. Consequently, nascent hydrogen is nothing else than ordinary hydrogen in thermic conditions or, speaking generally, in different physical conditions. To my mind, the expression nascent hydrogen is synonymous with hydrogen + calories. In fact, all the reactions produced with nascent hydrogen can be obtained quite as well with ordinary hydrogen and a high temperature; and the differences observed between the hydrogen resulting from different chemical reactions are simply due to the fact that these reactions do not develop the same quantity of calories.

Now is any further proof wished for that nascent hydrogen is nothing but $\text{H} + \text{cal}$. I take, for example, the action of hydrogen on sulphur. This body, as everybody knows, does not combine at the ordinary temperature with hydrogen, but the reaction takes place if that gas is in what is called a nascent state. On the other hand, M. Janario, by causing a current of hydrogen to pass over the melted sulphur has been able to determine a reaction between these two bodies. Is it not the same cause which lead

to the formation of sulphuretted hydrogen? In these two reactions the sulphur and hydrogen require a certain quantity of heat in order to combine; only in the first case that heat is provided by the combustion of lighting-gas, and in the second case by a chemical reaction. The physical conditions alone are changed, the hydrogen remains the same.

Dr. Phipson says that in 1858 he presented to the Society of Sciences at Haarlem, a memoir, entitled "Catalytic Force, or Study on the Phenomena of Contact," and in which he propounded a theory to explain the nascent state of bodies. I deeply regret that I was unacquainted with Dr. Phipson's interesting memoir, otherwise I should certainly have spoken of it in my notes. I must, however, say that the Chemical Dictionaries of Watts, Wurtz, and Selmi (on the article "Nascent Hydrogen"), and the principal chemical treatises do not mention in any way the experiments or theories put forth by Dr. Phipson; nor do they quote any chemist who took up the question before myself. It is right to say that M. Ste.-Claire Deville announced the existence of hydrogen in the nascent state, but he has not explained why that gas has greater affinity in the nascent than in the ordinary state.

I hope that when Dr. Phipson has read this paper he may perhaps accept my thermic theory of the nascent state of hydrogen. If, on the contrary, he fails to do so, I shall always be very happy to reply to any objection he may raise on the subject.

COMPARATIVE RESULTS OBTAINED WITH PREVIOUS ELECTRICAL OZONISERS, WITH DESCRIPTION OF A MODIFIED AND POWERFUL FORM.

By ALBERT R. LEEDS, Ph.D.

I.

In pursuing this enquiry, two objects were kept prominently in view:—1st. To obtain ozone by a method which could be relied upon at all times, and which would not only be independent of changes in temperature, humidity, &c., but require little or no supervision from the experimenter after once being set into operation. 2nd. To be able both to ozonise to a maximum the air or oxygen employed, and to employ large volumes of the ozonised gas.

The considerations summarised under the first head, finally caused the abandonment of a large Holtz machine, with which the experiments had been originally instituted. For, granting that by the use of suitable precautions the disturbing influences of atmospheric variations are eliminated, there still remains the necessity of a considerable amount of mechanical power to drive the machine.

With regard to the second point, we were assisted chiefly by Dr. Siemens's theoretical discussion of the principles involved in the construction of the induction tube which bears his name. In the experiments of Von Babo and Claus, with an ozoniser devised by the first-named, a very small volume of oxygen confined within the apparatus was subjected for many hours to the action of electricity generated by a powerful coil.* The maximum degree of ozonation was reached under these circumstances when the confined gas ceased to contract, and this maximum in various trials corresponded to 3·1 to 5·74 per cent of ozone. Brodie employed a Siemens tube, the interior and exterior surfaces being cooled to 0° or even -10° by ice or a mixture of ice and salt.† The highest percentage of ozone obtained in this manner was 6·5 per cent. A very powerful coil was employed, the

passage of the gas in each experiment occupying about thirty minutes, and the volume of the oxygen varying from about 100 c.c. to less than 300 c.c. Experiments were made with the view of ascertaining the effect of repeated electrification of the gas on passing it several times through the induction tube. It was found that on its fourth or fifth passage it contained no higher percentage of ozone than it did at the end of the first. These experiments agreed, therefore, with those of Von Babo and Claus, in so far as they likewise appeared to show that there was a fixed limit prescribed by the conditions of the experiment, beyond which the ozonation cannot pass, but differed in the respect that this limit was reached at once, while in those of Von Babo and Claus it was not attained until after powerful electrification for many hours.

But the object in the present enquiry was not to ozonise to a maximum a small volume of confined gas with the view of studying the properties of the gas itself. It was rather to ozonise to a maximum a *current* of air or oxygen, with a view of studying the reactions produced by it in the various substances with which it was brought in contact. It was also desirable, if possible, to employ a large volume of oxygen, and to keep it flowing rapidly, so that the total amount of ozone produced might be considerable. With these ends in view, the various forms of ozonisers hitherto proposed were tried, and failed to give, in our hands, the character of results sought for, until eventually, after many modifications, a development of the Siemens induction tube was arrived at, which has proven satisfactory.

After discarding the Holtz machine an induction coil of ten Bunsen elements (set up as two large cells) was employed. The axis of the wheel which operated the hammer used in making and breaking connections was also the axis of a pulley, connected by a belt with another pulley, run by a small electro-magnetic machine. This latter was driven by two additional Bunsen cells, and the speed so regulated that about 60 sparks would pass between the terminals of the coil per minute. As thus arranged, the coil was used to supply the electricity employed in making the trials on the various ozonisers employed. The results given are the best which we could obtain, and as such are put on record, without attempting to explain why, in some instances, they are so low.

WRIGHT'S OZONISER.*

The first tried was that of Prof. Wright, who, however, had made use in his own experiments of a Holtz machine as the source of electricity. The original description was followed closely. A straight glass tube, 20 c.m. in length and 2·5 c.m. in diameter, was closed at each end with paraffined corks, through the axis of which stout copper wires were passed, one carrying a terminal in the form of a ball, the other a terminal disc. Inlet and exit tubes were also let into the corks eccentrically to provide for the flow of gas.

A very slow stream of perfectly dry oxygen was made to flow through the apparatus. The maximum result was 0·76 m.grm. of ozone per litre. The oxygen used contained about 2 per cent of nitrogen. It was dried in this and the subsequent experiments by passage through a series of sulphuric acid dryers.†

HOUSSEAU'S OZONISER.

According to M. Houzeau's description (*Compt. Rend.*, lxxiv., 256), this is constructed of a straight tube, in the interior of which is a wire of copper, lead, or, better, of platinum, 40 to 60 c.m. long, passing through the upper

* *Am. Journ. Sci.*, [3], iv., 29.

† *Norm.*—Gianetti and Volta (*Berichte der Deutschen Chemischen Gesellschaft*, vii., p. 1462) with a similar apparatus, employing the Holtz machine, obtained 15 m.grm. of ozone per litre. Later (*ibidem*, ix., p. 84) they employed a form of Siemens induction tube, with the inner and outer tubes 2 c.m. and 2½ c.m. respectively, in connection with a Holtz machine. They found that the slower the stream and the lower the temperature, the larger the percentage. Between 5° and 10°, ½ litre of oxygen flowing per hour, they obtained 40 m.grm. ozone per litre. On substituting a Ruhmkorff coil, they obtained, under like circumstances, only one-third this amount.

* "Researches on the Constitution of Ozone," *Am. der Chem. und Pharm.*, xxi., p. 248.

† Sir B. C. Brodie "On the Action of Electricity on Gases," *Phil. Trans.*, 1872, p. 435.

side. A wire is wrapped around the exterior, which is a little longer than the interior wire, and connects with the other pole of the induction coil. With such an apparatus, and with a coil giving 2 to 3 c.m. sparks, Houzeau states that he readily obtained 60 to 120 m.grms. of ozone per litre of ozonised air, according as the experiment was performed at $+15^{\circ}$ or -30° . In one case he obtained 180 m.grms., and expressed the opinion that the complete conversion of oxygen into ozone was to be regarded as possible.

Two ozonisers of this form were made. The first, of a tube 55 c.m. long and 15 m.m. diameter. A spiral of platinum wire, 90 c.m. long, coiled around the middle portion of the exterior of the tube, formed one pole; a straight platinum wire passing along the axis of the interior of the tube, the other.

Results.—Oxygen used, a slow current flowing, 3.25 litres; ozone obtained, 16.07 m.grms., or 4.95 m.grms. per litre. Temperature, 20° . Second trial.—Oxygen used, 5.125 litres; ozone obtained, 19.85 m.grms., or 3.9 m.grms. per litre. Temp., 21° .

These results being highly disappointing, a more elaborate ozoniser of this class was constructed. The outside tube was 60 c.m. long and 18 m.m. diameter. Around the middle portion of this exterior tube, for a length of 40 c.m., was coiled a platinum spiral 177 c.m. long, with 32 turns. In the axis of this tube another glass tube 6 m.m. in diameter was placed, with a similar platinum spiral, but having 70 turns, and coiled in the opposite direction. The inner tube was kept concentric with the outer by two rings of glass beads.

Results.—Placed in a mixture of ice and salt. Oxygen used, $4\frac{1}{2}$ litres; ozone obtained, 19.03 m.grms., or 4.6 m.grms. per litre. No better result on second trial. Repeated.—Temp., 22° ; spark-length, 30 m.m. Oxygen used, 1.6 litres; ozone obtained, 3.78 m.grms., or 2.4 m.grms. per litre.

Since, in these and subsequent trials, Houzeau's ozoniser, in my hands, yielded equally unsatisfactory results, its use was abandoned.

BOILLOT'S OZONISER.

Its peculiar features are—1st. The use of gas-carbon as the material constituting the two poles; 2nd. The ozonised oxygen is allowed to flow through, and in contact with, one of these poles, instead of isolating the gas-carbon in a glass tube. The apparatus was constructed so as to embody the ideas of its author, as far as we understood them.*

The oxygen was conveyed through a glass tube, 60 c.m. long and 15 m.m. in diameter. This was closed with paraffined corks at both ends, through which smaller glass tubes were passed for inlet and outflow of gas. It was filled with coarse fragments of gas-carbon, each about the size of a pea. These were connected with one pole by a copper wire passing through the cork. The middle of this tube was surrounded by another tube, 19 m.m. in diameter, the space between them being filled with pulverised gas-carbon, and connected by a copper wire with the other pole.

Results.—Length of spark, 60 m.m. at beginning, 50 m.m. at close of experiment; temp., 20° ; time, two hours; amount of oxygen used, $4\frac{1}{2}$ litres; of ozone obtained, 11.91 m.grms. This is equivalent to 2.77 m.grms., or 1.29 c.c. per litre.

VON BABO'S OZONISER.†

The instrument of this description, as constructed by ourselves, was made as follows. 14 small thin tubes, 2.5 m.m. wide and 900 m.m. long, were joined together two by two, and thrust into a large tube, 21 m.m. inside diameter and 975 m.m. long. The thin tubes were kept

apart by means of little pieces of tubing, about 10 m.m. long and just large enough to fit over the other tubes. In each tube was placed a copper wire running through nearly its entire length, and connected with a platinum wire, and then they were sealed by the gas flame. The tubes in couples were so arranged that the platinum wire of one projected from one end of the large tube, and the platinum wire of the other projected from the other end. The bundles of wires were connected at each end with copper wires going off to the coil, the junction being made outside of the enveloping tube, which was provided with inlets and outlets for the gas.

Result.—Spark-length, 70 m.m.; time, thirty-five minutes. Oxygen used, 1 litre; ozone obtained, 9.15 m.grms. In the dark it was found that the discharge was not altogether silent, a few points of light being visible.

Thinking that a better result might be obtained by spreading the charge over a lesser extent, another ozoniser of this pattern was fitted up with only 5 pairs of tubes, all the other arrangements being the same.

Result.—Time, seventy minutes. Oxygen used, 1 litre; ozone obtained, 2.5 m.grms. Minute points of light were also noticeable in the dark along the sides of some of these tubes.

After these trials, the attempt to employ ozonisers of Von Babo's form was abandoned.

SIEMENS'S OZONISER.

The original description of this ozoniser, which was designed and constructed according to certain theoretical conceptions by W. Siemens (*Ann. der Phys. und Chem.*, cii., 120) more than twenty years ago, is as follows:—It is made of two tubes of the thinnest suitable glass. One of them, which is closed and the narrower of the two, is set inside the other in such a way that the annular space between them is of uniform width throughout. The outer is melted to the inner tube at one end, and at the other is narrowed down into an exit tube. The inlet tube is joined to the other end of the outer tube. The inner and outer surfaces of the glass tubes are coated with metallic foil and oppositely electrified.

With the objects in view, which were set forth at the beginning of this article, a very large number of ozonisers of this description were made, and the successive improvements, as shown by the increase in the amount of ozone obtained, rapidly led to the modification, which was finally adopted.

The first was constructed of a tube 50 c.m. in length and 35 m.m. in diameter, narrowed down at each end so as to unite with a straight tube passing through its middle, 17 m.m. in diameter. It is not necessary to fuse the tubes, so as to close the annular space, a joint of sealing-wax remaining unaffected. The inside of this second tube was covered with tin-foil, and formed one pole; the exterior of the outer tube was covered with 378 c.m. of tin-foil, and constituted the other pole. The lateral inlet and exit tubes for the passage of the gas were closed with small wash-bottles made of test-tubes, and containing sulphuric acid. This simple arrangement prevents backward diffusion of the ozone, and consequent destruction of the rubber connections.

Result.—Oxygen employed, 1 litre; ozone obtained, 7.18 m.grms.

After the experiments given in the present article were concluded, a Siemens ozoniser of the pattern known as Geissler's, was received from Bonn. The interior tube was 18 m.m., the exterior tube 27 m.m. in diameter, and the exterior surface of the latter over a length of 30 c.m. was coated with foil; about the same length of foil was contained in the inner tube. The outer tube was again surrounded by a final enveloping glass tube. All the joints and connections in this difficult piece of glass-blowing were of glass. In our own trials we were compelled to make the joints of sealing-wax. As this, however, lowered all the quantities found, their relative values and significance would still remain.

* *Comptes Rendus*, lxxv., 214 and 1712; *CHEMICAL NEWS*, xlv., 312; *Journ. Chemical Society*, N.S., x., 879.

† "Volumetric Relations of Ozone," *Ann. der Chem. und Pharm.*, Suppl. II., p. 297.

Result.—Spark-length, 31 m.m.; time, forty minutes. Oxygen used, 1 litre; ozone obtained, 28.16 m.grms.

A series of experiments was now instituted to determine the influence of the extent of electrified surface, its separating distance, duration of exposure of the current of oxygen to the silent discharge, and of the spark-length upon the amount of ozone obtained. With these objects we employed:—

I. Oxoniser of one metre in length, of which 850 m.m. were covered with tin-foil. Diameter of outer tube, 25 m.m.; width of annular space intervening, 1 m.m. Both inside and outside tubes of very thin glass.

Result.—Oxygen employed, 1 litre; ozone obtained, 23.65 m.grms. Spark-length, 85 m.m.; temp., 22°; duration of experiment, ten minutes.

II. Length of tube, 1 metre; length of exterior of outer tube covered by tin-foil, 925 m.m.; diameter of outside tube, 47 m.m.; diameter of inside tube, 25 m.m.; width of the annular space intervening, 11 m.m.; walls of the glass tubes thicker than the preceding.

Result.—Oxygen employed, 1 litre; ozone obtained, 5.8 m.grms.; time, ten minutes; temp., 22°; spark-length, 60 m.m.

Therefore, in the construction of tubes subsequently made, glass tubes as thin as could be obtained, and with as small an annular space between them as possible, were employed.

III. Using the same (II.) oxoniser, an experiment was made to determine the effect of prolonged exposure to the silent discharge upon the degree of ozonation. After filling with oxygen, the coil was thrown into operation for fifteen minutes, and then, the coil being still active, a current started, and allowed to flow until 1 litre of oxygen had passed through the oxoniser, the time required being ten minutes.

Result.—Oxygen employed, 1 litre; ozone obtained, 10.2 m.grms.; time, 15 + 10 = 25 minutes; temp., 22°; spark-length, 55 m.m.

IV. A similar experiment was tried with the first (I.) oxoniser.

Result.—Oxygen employed, 1 litre; ozone obtained, 21.54 m.grms.; time, 15 + 10 = 25 minutes; temp., 22°; spark-length, 55 m.m.

In this experiment a lower result in the total amount of ozone obtained is very noticeable as compared with I., in which, however, the spark-length was 85 m.m.

V. Using, therefore, the same oxoniser (I.) with the spark-length of .55 m.m., the experiment was repeated without setting the coil into action prior to beginning of the flow.

Result.—Oxygen employed, 1 litre; ozone obtained, 14.93 m.grms.; time, ten minutes; temp., 22°; spark-length, 55 m.m.

The foregoing experiments point out the proper directions in which to modify the apparatus in order, under given circumstances, to bring the degree of ozonation to a maximum:—

- (a.) To diminish the distance between the electrified surfaces, both by diminishing the thickness of the glass and by lessening the annular space between the two tubes.
- (b.) By prolonging the interval during which the oxygen is subjected to the silent discharge. There are evidently two ways of doing this—either by lessening the rate of flow, or, the intensity of static charge being preserved the same upon each unit of electrified surface, by increasing the length of the tubes. The latter method was adopted, since the principal object in view was that of obtaining the largest possible total quantity of ozone.
- (c.) Along with the increase in the length of the tubes, to raise correspondingly the length and number of the sparks (within the limits of the silent discharge).
- (d.) Since no notable elevation of temperature was perceived, in the case of large tubes operated in this

manner, it was deemed important not to complicate the apparatus by arrangements for depressing the temperature.

Since considerable time was required to line the inside of the inner tubes with tin-foil, some trials were made to learn whether a simpler arrangement would answer.

VI. The outside of the inner tube, this time of heavy glass, was coated with tin-foil, and the outside of the outer tube with the same, the length covered by the tin-foil being 900 m.m., and the annular space between the tubes 1 m.m.

Result.—Spark-length, 62 m.m.; time, 10 min.; temp., 21°; oxygen used, 1 litre; ozone obtained, 18.9.

VII. The tin-foil was transferred from the outside to the inside of the inner tube, everything else remaining the same.

Result.—Oxygen used, 1 litre; ozone obtained, 10.58 m.grms.

These trials demonstrated the importance of separating the electrified surfaces by glass walls. The tin was not noticeably affected, and the diminution of ozonation was probably due to the loss of tension consequent on removing one of the insulating surfaces.

VIII. To study the effect of prolonging the interval during which a certain volume of oxygen was subjected to ozonation, the oxoniser used in VII. was charged for fifteen minutes, the gas not flowing. The volume of oxygen contained in the annular space and ozonised in this manner was 70 c.c.

Result.—Spark-length, 55 m.m.; time, 15 minutes; oxygen used, 70 c.c.; ozone obtained, 2.31 m.grms., corresponding to 33 m.grms. per litre.

Since an increase in the length of the tube would also have the effect of prolonging the duration of ozonation, the foregoing experiment suggested those which follow, and ultimately led to a satisfactory solution of the problem in hand.

IX. A thin glass tube, 25 m.m. wide and 1 metre long, was coated with tin-foil over 815 m.m. of its outside surface. An inner tube was similarly coated on the inside over 815 m.m. Its inner coating was connected with copper-wire with one pole. Both ends of each tube were sealed with paraffin corks, the outer tube enclosing the inner, and the inner separated by glass rings by an annular interval of 1 m.m. from the outer. A duplicate ozonising tube was prepared, and the outer surfaces of the two connected together and with one pole of the coil, and likewise their inner surfaces with the other pole.

Results.—Spark length, 50 m.m.; temp. 21.5° C.; time, 30 mins.; oxygen used, 1 litre; ozone obtained, 52.54 m.grms.

Repeated, but the oxoniser charged for 5 minutes before beginning the flow of oxygen; ozone obtained, 57.83 m.grms.

Repeated, the charging being begun 15 minutes before flow; ozone obtained, 71.82 m.grms.

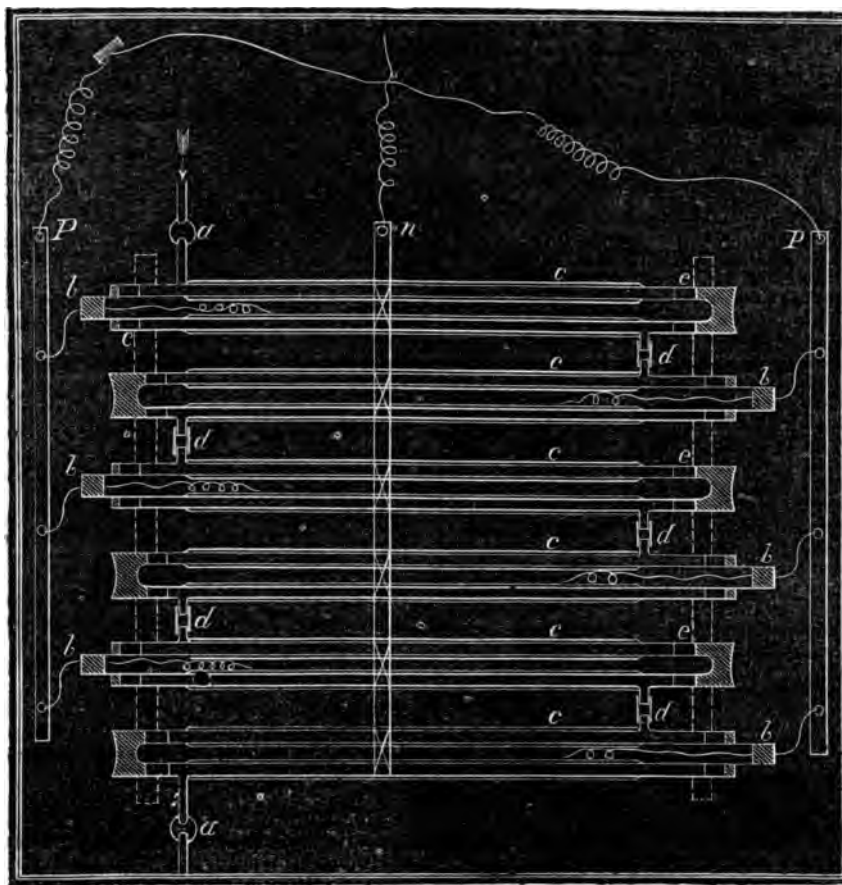
Finding, as the result of other variations in the lengths and proportions of the tube, the conclusions arrived at above were confirmed, the form of oxoniser as finally used and adopted was constructed, and will be found figured in the text. Each induction-tube is a modified Siemens tube, and may be called a Siemens's element. Six of these elements are connected together and supported on a frame, constituting what we may term an ozonising battery. One of these frames is fitted on above another, the end elements of the two batteries being suitably connected, and in this way, by repetition of similar parts, an ozonising tube of as great length as desired can be made and handled without inconvenience. Or the electrical charge can be divided between several ozonising batteries and the supply of oxygen as well, so that a number of currents of ozonised gas can be used at the same time.

Each element is made of a tube of thin hard glass, 60 c.m. long and 21 m.m. inside diameter, with the inlet and outlet tubes 6 c.m. from each extremity. The space between these two latter tubes is coated with tin-foil. The inside tube is a little longer, one end being rounded; the

other, after the interior has been coated with tin-foil, is closed with a dry cork, through which the copper connecting wire passes. The space between the rounded end of the inner tube and the outer tube is nearly filled by a ring of glass cut from a tube of suitable bore, and the space closed by dipping in molten sealing-wax. In coupling the elements together, the exit-tube of the first is joined to the inlet tube of the second by a wrapping of strips of muslin, which are bound by flower-wire, and made gas-tight by a coating of molten paraffin applied with a brush. The first inlet tube of a battery and the last exit tube are made parts of small sulphuric acid wash-bottles, by means of which the rubber or kerite connections with the other parts of the apparatus are protected from backward diffusion of the ozonised gas on the one hand, and a convenient attachment is made upon the other.

the work required, it was used almost daily for three months, the average yield of ozone, with the battery in good condition, being 72 m.grms. per litre, or about 5 per cent.

To obtain the best results with such a battery the following precautions should be observed:—(1) The connections at *dd* should preferably be ground, and the other joints be made by fusion. (2) The number of elements used should be proportioned to the strength of the coil, the maximum being obtained when the whole interior is luminous, but without sparks, in a darkened room. (3) The feeble inductorium should be replaced by one giving a large number as well as a great length of sparks. (4) The exterior foil should be covered with an outer enveloping tube of glass. (5) The temperature of the ozonising battery should be prevented from rising by



DESCRIPTION.

aa, small sulphuric acid wash-bottles. *bb*, corks closed by melted sealing-wax. *cc*, outside coating of tin-foil. *dd*, connections of paraffined cotton-cloth. *ee*, rings of glass. *pp*, copper strips connecting with inner coating and one pole. *w*, copper strip fastened to all the outer coatings and the other pole.

Results.—With a battery of 3 elements—Time, 30 mins.; oxygen used, 1 litre; ozone obtained, 22.8 m.grms.

With a battery of 6 elements—Time, 30 mins.; oxygen used, 1 litre; ozone obtained, 51.74 m.grms.

Repeated, after an interval, but with same number of elements—Oxygen used, 1 litre; ozone obtained, 49.5 m.grms.

With two batteries, 12 elements in all—Spark-length, 45 in.m.; time, 30 mins.; temp. 24°; oxygen used, 1 litre; ozone obtained, 69.93 m.grms.

In this shape, the apparatus proving adequate to perform

placing it within a refrigerating chamber, and surrounding it by dry air kept at 0°.

Examination of Commercial Gums.—E. Masing.—The author compares the behaviour of the Arabian, Asiatic, American, and Australian gums with reagents, and concludes that the Arabic acid, present in all, occurs in modifications influenced by the presence of the mineral constituents.—*Arch. Pharm.*, xii., 216.

ACTION OF LIME ON SILICA IN MORTAR.

By W. B. ROBERTS, M.S.A.

HAVING found in the recent analysis of some specimens of old mortar from the walls of a building, erected about two hundred years ago, considerable traces of hydrated silica, it occurred to me that possibly the hardening or setting of mortar might be due to some chemical action occurring between the lime and the silica when these ingredients were mixed, whereby some proportion of the silica was caused to assume the gelatinous form; that this being then incorporated by the usual mixing process, subsequently solidified, binding the whole bulk with a hard network of silica. To test this, I obtained two good specimens, one from the exterior, and one from the centre of a wall which was two feet thick, for the purpose of making a careful analysis of each. In order to compare the results with those of some experiments presently to be described, I also examined a sample of mortar from a building of comparatively recent date.

Subjected to the mechanical test of a gradually increasing pressure, the two older samples proved about equal in hardness, while both of them were harder than the third. The samples taken for analysis were quite free from brick, and few of the grains of sand were of greater weight than about 1 grm.

The specimens were crushed fine in a steel mortar, and 200 grms. of each treated for twelve hours with sulphuric acid (sp. gr. 1.65); the temperature was then gradually raised, until the excess of acid was completely expelled; after cooling the residue was boiled up repeatedly with water; the insoluble residue gave the total silica—conveniently referred to as silica, silica free, and sand.

To find amount of soluble silica, the insoluble residue was boiled with a saturated solution of sodium carbonate for a considerable time; the liquid, after filtration, was treated with hydrochloric acid and evaporated to dryness in the usual way, and gave the amount of free and combined soluble silica in the mortar.* Then, to ascertain how much, if any, free hydrated silica, i.e., silica which was or had been gelatinised, existed in the mortars, 200 grms. of each were treated with saturated solution of sodium carbonate exactly in the manner just described for separating the soluble silica from the sand. The percentage of previously gelatinised silica was confirmed by treatment with pure hydrofluoric acid. The other constituents were estimated by the ordinary methods. The following were the results obtained:—

	I. From Exterior of old Wall.	II. Interior of old Wall.	III. More recent Mortar. Exterior.
Sand	67.12	67.19	65.54
Combined silica ..	1.12	0.93	0.18
Free hydrated silica ..	0.68	0.40	0.09
	68.92	68.52	65.81
Calcium carbonate ..	25.05	28.04	31.84
Magnesium carbonate	0.68	0.60	0.09
Sodium carbonate ..	0.90	1.41	trace
Lime, originally as silicate	1.05	—	—
Soda	0.41	—	—
Calcium sulphate ..	0.49	0.43	0.19
Sodium chloride ..	0.17	0.14	0.08
Oxide of iron, alumina	0.02	0.02	0.01
Water, hygroscopic ..	2.09	1.02	1.78
	99.78	100.18	99.80
Carbonic anhydride ..	11.78	13.18	15.033

To find as far as possible whether the original lime had contained any calcium silicate—this being, as is well

known, produced sometimes in comparatively large quantities during the burning of different limestones—as much of the lime as could be picked out free from sand by the aid of a lens was tested for free or combined soluble silica in the same way as the mortar. The results were:—

	I.	II.	III.
Soluble silica per cent ..	0.17	0.20	0.11

Further, to test whether, in a comparatively short time, the sand would be attacked by freshly slaked lime, 300 grms. of fine sharp sand, free from soluble silica, were placed in each of three bottles under the following conditions for six months, after which time analysis gave the results stated below:—

	Soluble Silica Per cent.
I. Mixed with 150 grms. pure lime, and sufficient water to make a thick paste, and prevented from absorbing CO ₂ ..	0.09
II. Same as I., but with the addition of 2 grms. of mixed carbonate of soda and potash	0.17
III. Same as II. with occasional slow current of CO ₂ , besides constant access of atmospheric air	0.08

From the general results of the above analysis and experiments, I conclude that the accepted theory of the hardening of mortar, namely, that it is due to the absorption of carbonic acid, is unshaken.

In the cases of the experiments I. and II. the conditions have been very favourable for the action of the lime upon the silica, but its effects are manifestly too small to materially influence the hardness of the resulting mortars. However, samples I. and II. were harder than III., although I. contained much less CO₂ than III. My general conclusion may be summarised as follows:—

- (1.) Practically no gelatinisation of silica occurs in the manufacture of mortar.
- (2.) That under the ordinary conditions of access of atmospheric air the lime in mortars becomes gradually dehydrated, absorbs carbonic acid, and forms neutral carbonate.
- (3.) That the absorption of carbonic acid is very slow.
- (4.) A slight action takes place between the lime and the silica, though this appears to be too small to determine its nature.
- (5.) That, although even the small proportion of dry silicates slightly increases the hardness of a mortar, the ordinarily sufficient hardness of mortar is obtained by simple dehydration and carbonation.

These conclusions appear to be confirmed by the fact that lime already containing a small proportion of carbonate is preferred to pure lime for making mortar.

University of London.—The following is a list of the candidates who have passed the recent Second B.Sc. Examination:—(First Division).—S. G. H. Barfield, private study; G. Brown, private study; G. J. Burns, private study; W. D. Halliburton, University College; S. L. Hart, St. John's College, Cambridge; R. S. Heath, Trinity College, Cambridge; G. B. Hoffmeister, Gonville and Caius College, Cambridge; E. Hopkinson, Emmanuel College, Cambridge; A. H. S. Lucas, private study; J. G. Ridesdale, Guy's Hospital and private study; G. T. Smith, Modern School, Bedford; J. W. W. Waghorn, private study; T. W. Williams, University College and Royal School of Mines. (Second Division).—J. A. B. Bennett, Downing College, Cambridge; W. Fawcett, private tuition; J. J. Fletcher, B.A. Syd., Royal School of Mines and University College; S. J. Hickson, Downing College, Cambridge; W. Overend, University College and St. Bartholomew's Hospital; T. Purdie, University of Würzburg and private study; E. H. Rennie, M.A. Syd., St. Mary's Hospital and private study.

* Thorpe, "Quantitative Analysis," p. 185.

PROCEEDINGS OF SOCIETIES.

NEWCASTLE CHEMICAL SOCIETY.

General Meeting, October 23, 1879.

Mr. R. C. CLAPHAM, President, in the Chair.

THE minutes of last meeting were read and confirmed.

The Treasurer's statement was presented.

The PRESIDENT—The next business is the reading of the Committee's Report. Members will learn with regret from the Report that Mr. Proctor has been compelled, from failing health, to resign the office of Secretary and Editor of the Transactions. Mr. Proctor has filled the office of Secretary very ably, and I can testify personally that he has brought an amount of knowledge and industry to bear on the reports which was highly commendable. I am sure the Society will agree with me when I say that we have lost in Mr. Proctor a most able and conscientious officer. We can only hope that his successor will endeavour to emulate him.

The Committee's Report was then read as follows:—

Committee's Report.

Your Committee beg to report to the twelfth annual meeting of members that the number now on the Society's books is 141, against 128 last year. The number who joined the first year of the Society's existence was 96, and it is therefore satisfactory to notice a gradual improvement in the numbers. Owing to indisposition and business engagements our much-esteemed colleague, Mr. B. S. Proctor, has been obliged to resign the office of senior Secretary, and the Committee have now to recommend that Mr. J. T. Dunn, of the College of Physical Science, should be elected in his place.

The papers read during the last session were as follows:—"Rosan's Process for Desilverising Lead," by Mr. N. Cookson; "On the Estimation of Nitrous and Nitric Acids," by Dr. Lunge and Mr. Maclear; "On Noxious Vapours," by Mr. Hill; "On a Spring Water containing Manganese," by Mr. H. S. Pattinson; "On Nitric Nitrogen in Guano," by Mr. Tatlock; "Chemical Jottings," by Mr. H. R. Procter; "The Smoke of an Electric Lamp," by Mr. B. S. Proctor.

At the conclusion of the reading of many of the above papers there were useful and interesting discussions, which were the means of adding much information not touched upon in the papers. This was especially the case in an important paper by Mr. Cookson, on the "Separation of Silver from Lead."

In conclusion, the Committee wish again to express a hope that there will be, during the present session, more contributions from the general body of the members, many of whom are so well able to contribute important scientific information.

Mr. SCHOLEFIELD—I move the adoption of the Report, and I should like at the same time to propose a vote of thanks to Mr. Proctor for the able manner in which he has fulfilled the duties of Secretary. With all the remarks which Mr. Clapham has made on this subject I heartily concur.

Mr. H. R. PROCTER—I beg to second the adoption of the Report, and also the vote of thanks to Mr. Proctor.

Both motions were then put to the meeting and carried unanimously.

Mr. William Martyn and Mr. C. E. Stuart were elected members of the Society.

The PRESIDENT—If no member has any remarks to make on the papers which come up for discussion to-night I will proceed to read the paper which I have prepared for this meeting. Although there is no particularly new matter in the paper, I thought it might be useful at this early stage of the production of steel from Cleveland iron

to enter the facts, as far as they are known, on our minutes; at any rate they might be historically interesting. I have found it rather difficult to say anything about a process which is as yet only in its infancy; but I have consulted a good many ironmasters on the main facts contained in the paper, and have at any rate their authority for its correctness.

"Extracts relating to the Manufacture of Steel from Cleveland Iron," by R. C. CLAPHAM, F.C.S. In an inaugural address, which I delivered last year, and towards the end of that paper, I referred to important experiments that were then being conducted with the object of producing steel from Cleveland pig-iron, in the large way. Since then some practical progress has been made, and steel is now being produced in an experimental plant erected for the purpose by a well-known iron firm at Middlesbrough—at any rate to the extent of a few hundred tons weekly, some quantity of which has already been sold as railway rails. The total quantity of Cleveland pig-iron made annually is about 2,000,000 tons. The importance, therefore, to the North of England of the ultimate success of the steel process cannot be exaggerated, even should it require, as it may possibly do, some years to accomplish.

A very valuable paper was read by Thomas and Gilchrist at the spring meeting of the Iron and Steel Institute held in London, "On the Elimination of Phosphorus in the Bessemer Converter." The difficulty experienced in separating the phosphorus in a puddling furnace, or in a Bessemer converter, is shown in this paper to arise from the hitherto almost universal practice of using fire-bricks or other silicious linings, the silicon when oxidised acting upon the materials as an acid; whereas an earthy or alkaline base would appear to be necessary both as a lining and as an addition to the melted pig, to give the phosphorus, at the moment of oxidation, a base with which to unite and separate itself from the metal. Mr. Richards has recently adopted a plan of blowing in fine lime through the tuyeres amongst the melted pig, with the object of bringing the base and the metal into more immediate contact. As a cheap material was absolutely requisite, and a material which would stand the great heat in the furnace, lime or magnesian limestone, which is found in such extensive deposits in Durham and Yorkshire, has been fixed upon for this purpose.

From the paper referred to I propose to make a few extracts as follows:—"An examination of the general conditions attending the removal of phosphorus in puddling and refining, taken in connection with the well-known action of silica on phosphate of iron at a high temperature, and the fact that in many other processes in which the temperature is very high, the elimination of phosphorus is not effected, seems to justify the belief, which has doubtless suggested itself to other members, that it is to the silicious (or fire-brick) lining of the ordinary converter and to the consequent necessarily silicious nature of the slag that the one defect of the Bessemer process is due. Under this conviction, at all events, experiments were commenced by the authors about three years ago on the effect of basic lining and basic additions in the several steel-making processes. The lining in these experiments consisted of limestone and silicate of soda, a mixture which had been found to answer well in earlier trials. Some fifty or more blows were made in a vertical converter, and the products analysed, and it was found that when using a basic lining it was generally necessary to continue the blow for over forty seconds after the flame dropped, in order to bring the phosphorus down very low. With this proviso the elimination of phosphorus could be secured with absolute certainty. With a silicious lining on the other hand, the retention of the phosphorus in the metal was as usual, equally invariable even when, as in Mr. Bell's experiments, the blow was continued till a considerable proportion of the iron was oxidised. At the same time more phosphorus and less silica would be found in the slag obtained under these conditions than appears

to be the case where large quantities of metal are treated under similar circumstances. It would appear that the presence of a considerable amount of lime is highly favourable, and on a large scale essential, to the removal of phosphorus. Phosphorus was not removed until the slag was sufficiently basic; the effect of large basic additions, in combination with a basic lining, was tried with the object not only of obtaining a highly basic slag at an early stage of the blow, but of rendering the operations independent of the wear of the lining, by which alone the basic character of the slag is otherwise obtained. Advantage was also taken of the fact that lime and oxide of iron are fusible in many proportions. The mixture generally used consisted roughly of one part by weight of 'Blue Billy' and two of lime; this melts in an iron crucible, and may be readily added in a molten condition. It was found, however, that by simply throwing into the converters cheap basic materials, such as lime, even without previous heating, before the pig was introduced, very satisfactory results were obtained without the disadvantage of over-blowing. The following example may be given:—

The pig used contained 1·19 per cent	} phosphorus.
1·39 " "	
And the steel produced contained 0·29 per cent	
0·13 " "	
0·04 " "	} phosphorus.

The amount of base which it is necessary to add with Cleveland pig generally exceeds 2 cwts. on the ton of iron, and the presence of an excess of base in the slag secures an essential condition of success."

Highly-fired magnesian limestone bricks for the lining are stated to have fulfilled the expectations which were entertained of their probable value. A patent has been taken out by Mr. E. Riley, F.C.S., for the manufacture of pressed limestone bricks. Various liquids were tried with the object of making the lime adhere under pressure while forming the bricks, and at last petroleum oil, coal oil, and resin oil were found to answer the purpose, used in the proportion of 5 to 10 parts of oil to 100 of lime.

The bricks are subject in a kiln whilst baking to a sharp, bright heat, during which operation a considerable shrinking in size takes place, and at the same time the petroleum or other oil slowly burns out, leaving a hard, compact brick, suitable for a lining. A limestone now used as a building stone in London has so far been tried by Mr. Riley in his experiments, which consists of nearly pure magnesian limestone, containing 98·94 per cent carbonates of magnesia and lime.

At the meeting of the same Society, held in Liverpool in September last, similar interest was shown in the practical separation of phosphorus. Three papers were read on the subject.

Mr. A. Pourcel stated that in the dephosphorisation of iron three essential elements must be taken into consideration in the early stage, even of the blast furnaces:—1st. The chemical composition of the cinder; 2nd. The temperature of the place in which the operations are performed; 3rd. The atmosphere in which the reactions take place. The same iron ores, he says, according to the temperature applied in the blast furnace, will produce grey, mottled, white, or cold spongy pig. He found the following results from experiments extended over a period of five years:—

Grey pig contains 0·60 per cent phosphorus on an average.
Mottled " 0·38 " " "
White " 0·30 " " "
Spongy " 0·18 " " "

The highest heat in the blast furnace producing grey pig, and consequently containing the most phosphorus.

Mr. R. Brown read a paper, in which, instead of getting rid of the phosphorus, he advised that it should be retained in the pig even up to 1·5 per cent, and that its effects should be what he styles "neutralised" by the addition of bichromate of potash in crystals in the pro-

portion of one-half part of chromate to 100 parts of pig in a molten condition. The iron so produced was said by the author to be equal to good steel, and suitable for steel castings, or steel plates for shipbuilding. This process, however, would appear on the face of it (even should further trials prove the conclusions to be correct) to be an expensive one, from the difficulty of providing bichromate of potash in sufficiently large quantities and at a cheap rate, owing to the comparative scarcity of the chrome ores.

A third paper, by Mr. H. B. BULL, suggested the separation of the phosphorus by converting it into phosphuretted hydrogen by means of steam blown into the mass of melted iron. However correct this process might be from a chemical point of view, it was shown by Dr. Siemens as likely to be impracticable from the probable rapid cooling of the iron during the time of blowing in the steam.

Possibly in another year we may have still further progress to report in the highly important operation of producing steel from Cleveland iron. In the meantime, large quantities of Bilbao iron ores and other ores suitable for making steel continue to be imported into this district, which appears to be sufficient proof that the new processes are not yet practically completed.

Mr. BERKLEY—The important part in the process appears to be the material of the bricks used for lining the converters. I had some talk with a gentleman who manufactures these bricks, and was informed that they were made from magnesian limestone, and require to be heated to a very high temperature in their manufacture. When the bricks are heated alone, they will stand any temperature; but if they come in contact with silica, or with the ordinary silicious bricks, they readily fuse. No doubt bringing magnesia into contact with the iron is the right thing. In this district the marl of the limestone from Fulwell is put into the puddling furnaces, and has a good effect on the iron if not used in excess. If too much is put in, however, it makes the iron red-short. I may say that I do not entertain the gloomy anticipations as to the future of Cleveland if these steel experiments should fail, which are held by some people. For many purposes iron will still be used, and steel will never supersede it. I think that steel rails, however, will always continue to be used; the steel at present exhibits great differences in quality, but doubtless as the manufacture improves these differences will disappear, and we shall be able to obtain steel of uniform quality, even in a large lot.

Mr. STUART—I should like to ask, firstly, are the bricks so highly heated in their manufacture as to drive off all the CO₂ from the lime and magnesia? Secondly, are they very refractory, as compared, for instance, with glauco or other silicious bricks? And thirdly, are they hard to handle? The high temperature at which revolvers are now worked renders it difficult to get bricks to stand the heat; we have to pay about 72s. per 1000 for bricks sufficiently refractory for revolver linings, while local bricks could be obtained for about 37s. per 1000. And hardness is a very desirable quality, for unless the bricks are capable of standing a good deal of knocking about, there is a great waste through breakage in the handling. If then these magnesia bricks are free from CO₂, refractory, hard, and obtainable at moderate cost, they would probably be very suitable to line revolver heating furnaces.

Mr. WATSON—I happened to be at Eston a few days ago, and saw the manufacture of the bricks. I could not, of course, say from their appearance whether they retained carbonic acid or not, but they are, at any rate, exceedingly hard, for I tried to break one and failed.

Mr. PROCTOR—Did the bricks exhibit signs of semi-fusion at all?

Mr. WATSON—Yes; they showed marks of fusion on the surface. I was told that they were burnt at a very high temperature, and lost about 50 per cent of their bulk in the process.

Mr. PROCTOR—Do they stand moisture? For if they

are simply lime and magnesia they might be expected to slack on exposure to air and damp.

Mr. STUART—Mr. Proctor's question is very important; possibly the semi-fused coating may have the effect of protecting them to some extent.

Mr. CLAPHAM—Lime formed into a very hard and compact mass is difficultly acted on even by acids.

Mr. SWAN—I should like to say that the idea of magnesia bricks is not new. Similar bricks were made in Paris some years ago by Tessié du Motay and Maxwell Lyte, of pure magnesia.

Dr. AFFLECK—Magnesia does not slack, and that is the chief reason why a magnesian limestone is sought for. I do not see why lime should not slack, however hard it may be.

Mr. BERKLEY—In making hematite pig, where the slag contains a very large amount of lime, we find that the slag does not stand the atmosphere at all but crumbles away, while our Cleveland slag is unaffected by it.

Mr. PROCTOR—How does lime which has been exposed to the oxyhydrogen flame behave?

Mr. SWAN—The point on which the flame impinges is so small that it would be difficult to separate from the rest of the lime, and I don't know that any observations have been made on it.

Mr. STUART—Hard-burnt lime is very difficult to slack, and very likely the same may be the case with these bricks. I should like to know the cost of the bricks.

Mr. BERKLEY—I believe 60s. per 1000.

Mr. WATSON—I noticed that the magnesia bricks at Eston were kept under cover, while the silicious bricks were exposed to air.

Mr. FRANCE then proposed, and Mr. WATSON seconded a motion, that a vote of thanks be given to the President for his interesting paper.

The motion was carried unanimously.

The meeting then terminated.

CORRESPONDENCE.

DETERMINATION OF SULPHUR IN COAL.

To the Editor of the Chemical News.

SIR,—The process for determination of sulphur in coal and coke by combustion with calcium hydrate, which was referred to by Dr. Divers in a note to a paper read at the Chemical Society's meeting, as reported in your last issue, did not owe its origin to Mr. Pattinson, but to myself, and was published by me in the CHEMICAL NEWS in 1874. I may also state that the method given in the paper just read before the Chemical Society was communicated to me by Mr. James Hargreaves, of Widnes, as far back as 1869.—I am, &c.,

W. F. K. STOCK.

5, Dixon Terrace, Darlington,
November 17, 1879.

MISCELLANEOUS.

Improvements in Gas-Furnaces.—We have frequently had occasion to call attention to the improvements effected in blowpipe apparatus and gas-furnaces by Mr. Fletcher, of Warrington. His improved blast-furnaces, crucible and muffle furnaces, automatic blower and aspirator, and chemical blowpipe are all great improvements on the old forms of apparatus. He has recently made a "Solid Flame Burner" and a "New Evaporating Burner," both of which we have used in our laboratory, and the results fully justify the advantages referred to by the inventor in his catalogue. The solid flame burner is five inches high, and requires much less gas than the old form of heating

burner to do the same amount of work. It is, moreover, free from smell. The flame appears to be the same temperature throughout. The inventor says that it will boil half a gallon of water in a flat copper kettle in five minutes, and will melt 6lbs. of lead or solder in an iron ladle in seven minutes. The evaporating burner for glass and porcelain vessels and general laboratory work is a great improvement on the ordinary coil burners in use, owing to the fact that no currents of cold air, which are so fatal to glass and porcelain dishes, can reach the vessel, as is the case with all coil burners. The flames are blue and smokeless, and are not liable to be extinguished with a splash, being raised above the body of the burner.

The Royal Society Medals.—The Council of the Royal Society have awarded the Copley Medal to Prof. Rudolph J. E. Clausius, of Bonn, for his well-known researches upon heat; the Davy Medal to M. Lecoq de Boisbaudran, for his discovery of gallium; a Royal Medal to Mr. William Henry Perkin, F.R.S., for his synthetical and other researches in organic chemistry; and a Royal Medal to Prof. Andrew Crombie Ramsay, F.R.S., for his long-continued and successful labours in geology and physical geography.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin
No. 12, 1879.

Methyl-pyrogalllic Acid and the Formation of Pittacall.—A. W. Hofmann.—The author has already shown that cedriret and pittacall are both derived from the secondary methyl-ether of pyrogalllic acid. He now regards pittacall as a base corresponding to rosanilin, a hexa-methoxylised rosolic acid. The author further describes the salts of eupittonic acid, the oxidation of a mixture of pyrogalllic acid diethyl-ether with methyl-pyrogalllic acid dimethyl-ether. The acid formed from a mixture of these two ethers in presence of ammonia readily passes into the corresponding tinctorial base.

Decomposition of Sulph-hydantoin by Barium Hydrate.—R. Andreasch.—The formation of thioglycolic acid from sulph-hydantoin by the action of an alkaline solution explains in the simplest manner why all attempts to desulphurise sulph-hydantoin in order thus to arrive at a glycolyl-amid lead merely to negative results.

Reaction with Iron Peculiar to Thioglycolic Acid.—R. Andreasch.—If the slightly acidulated solution of a thioglycolate, or also a solution of the free acid, is mixed with a drop of a very dilute solution of ferric chloride, a very transient indigo blue colouration appears in the liquid, but if the liquid is then made alkaline with a few drops of ammonia a dark red colouring, bordering on violet, appears, and becomes more intense on shaking with air. If too much ferric chloride is added, on supersaturation with ammonia hydrated ferric oxide falls without any change of colour.

Behaviour of Hæmatoxylin on Dry Distillation.—R. Meyer.—From the distillate a crystalline phenol mixture is easily separated, in which pyrogalllic acid (as gallein and resorcin; as fluorescein) are readily detected. Hæmatoxylin is a mixture of the above mentioned phenols.

Nitronaphthoeic Acids.—A. G. Ekstrand.—Not adapted for useful abstraction.

Determinations of Specific Gravities.—F. W. Clarke.—Not suitable for abstraction.

Amido Derivatives of Diphenyl-amin.—R. Nietzki and O. N. Witt.—An account of mono- and di-amido-diphenyl-amin.

MEETINGS FOR THE WEEK.

SATURDAY, 22nd.—Physical, 3. "On a Retention Image Photometer," Dr. F. Guthrie. "A Hint as to the Constitution of Chlorine," Prof. A. W. Rucker. "The Influence of Heat upon Certain Forms of Induction Coils," R. C. Shettle, M.D. "The Crossley Telephone Transmitter," Walter Emmott.

MONDAY, 24th.—Geographical, 8.30.
Medical, 8.
Society of Arts, 8. (Cantor Lecture). "Chemistry of Bread and Bread Making," Dr. Graham.

TUESDAY, 25th.—Civil Engineers, 8.
Anthropological Institute, 8.

WEDNESDAY, 26th.—Society of Arts, 8.

THURSDAY, 27th.—Royal, 8.30.
Philosophical Club, 6.30.

FRIDAY, 28th.—Quekett Club, 8.

CHRISTMAS LECTURES.

ROYAL INSTITUTION OF GREAT BRITAIN.

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TO INVENTORS AND PATENTEES.

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AND
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THE CHEMICAL NEWS.

VOL. XL. No. 1044.
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THE PERIODIC LAW OF THE CHEMICAL ELEMENTS.

By D. MENDELEEF.

(Continued from page 244.)

I. PRINCIPLES OF THE PERIODIC LAW.

ANALOGIES have long been noticed between the elements of high atomic weights and those of low ones. Claus observed that Os, Ir, and Pt, whose atomic weights are about 195, have analogous properties with Ru, Rh, and Pd, which have much lower atomic weights, viz., about 105. Marignac noticed the analogy of Ta=182 and W=184, with Nb=94 and Mo=96.

To Au and Hg correspond the lighter analogues Ag and Cd, as well as the still lighter ones Cu and Zn.

Cæsium and barium are analogues of potassium and calcium, and so on.

Comparisons of this nature lead to the idea of classifying all the elements according to their atomic weights, and in making this list we find reciprocal relations of a remarkable simplicity. As an example we will give all the elements whose atomic weights are between 7 and 36. These atomic weights are here arranged in arithmetical order according to their value.

Li = 7; Be = 9.4; B = 11; C = 12; N = 14; O = 16; F = 19;
Na = 23; Mg = 24; Al = 27.3; Si = 28; P = 31; S = 32; Cl = 35.5.

It is plainly shown that the character of the elements is subjected to regular modifications, and little by little, step by step, as the atomic weights vary, it is also periodically modified; that is to say, in the same manner in the two series, so that the corresponding members of these series are analogous. Na and Li, Mg and Be, C and Si, O and S, and so on. The corresponding elements in the two series have the same kind of compounds, and they have, as it is generally said, the same atomicity. The most important circumstance is that the forms of compounds of intermediate elements obey the laws which have been discovered in comparing the hydrated and oxygenised compounds of the elements which precede and follow after. This regularity proves that the elements just mentioned form two natural series in the order in which they are placed, and can neither of them have any intermediate members. Thus the last four members alone can combine with hydrogen, forming, if R represents an element:—

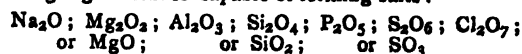


The constancy or destructibility of these hydrogen compounds under various influences, as well as their acid character or the faculty of changing hydrogen for metals, and other properties, are gradually modified, according to the position of the elements in the series. HCl is a very decided acid of great stability; H₂S is a weak acid, decomposed by heat; in H₃P the acid character has completely disappeared, and the instability has increased; again, it is much more decided in H₄Si.

All the elements of the second series enter into combination with oxygen. It is therefore in these compounds that we can best observe how the properties of elements are gradually modified, according as the atomic weight changes. As examples for comparison of this kind we will take the higher anhydrous oxides, by which I mean those which correspond to water, which can combine with it and form hydrates, and unite amongst themselves to form salts. We will not here take notice of those oxides which by their composition and properties correspond to

peroxide of hydrogen (such as Na₂O₂), because there are only a small number of elements capable of forming such compounds.

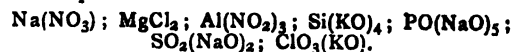
The seven elements of the second series give the following higher oxides capable of forming salts:—



Thus, in a determinate order, seven forms of oxidation known to chemists correspond to the seven members of the above series. Although they have long been discovered, their connection with the fundamental properties of elements remains unknown. We see, by examining these seven forms, that to their position corresponds a diminution of the basic properties and an increase of the acid properties. They give the following normal saline derivatives:—



For example—



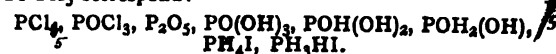
We can represent X by the following elements:—



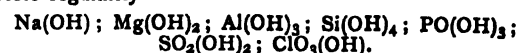
And, further—



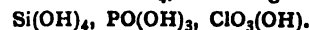
To PX₅ correspond:—



In the composition of hydrates we again notice a complete regularity—



In the hydrates there are only four hydroxyls, (OH)₄; and in the same manner in the oxides there are only four oxygen atoms, O₄. The highest forms of combination known with an element R are RO₄, RH₄, and R(OH)₄. No stable hydrate, S(OH)₆, corresponds with the type SX₆, although corresponding basic salts are known. This hydrate is transformed by elimination from 2H₂O into SO₂(HO)₂, which contains O₄, resembling in this manner,



It is not only in the forms of compounds of elements placed according to the amount of the atomic weights that we notice a regular connection, but also in their other chemical and physical properties.

At the beginning of the series are the bodies of a decided metallic character; at the end are the representatives of the metalloids. The former possess basic properties, the latter acid ones; while the bodies in the middle have intermediate properties. In Li₂O and Na₂O the basic properties are more clearly defined than in BeO and MgO; in B₂O₃ and Al₂O₃ they are only feebly shown, and these compounds have some acid properties. CO₂ and SiO₂ have only acid properties, although in only a small degree. This character is more pronounced in N₂O₅ and P₂O₅; and again in SO₃ and Cl₂O₇.

To show with what regularity the physical properties are modified in the above-mentioned series we give the modifications of the specific weights and the atomic volumes of the members of the second series:—

	Na.	Mg.	Al.	Si.	P.	S.	Cl.
Density	0.97	1.75	2.67	2.49	1.84	2.06	1.33
Atomic volume	24	14	10	11	16	16	27
	Na ₂ O.	Mg ₂ O ₂ .	Al ₂ O ₃ .	Si ₂ O ₄ .	P ₂ O ₅ .	S ₂ O ₆ .	Cl ₂ O ₇ .
Density	2.8	3.7	4.0	2.6	2.7	1.9	?
Atomic volume	22	22	25	45	55	82	?

The volatility diminishes in the members of the commencement of the first series from Na to Si; from thence

it increases. The same observation applies to the oxides, amongst which the middle ones, MgO , Al_2O_3 , SiO_2 , are the most refractory.

The compounds of metals at the beginning of the series with other metals are called alloys; they have the look and the properties of metals. Even in the case of phosphorus and sulphur these properties are not completely effaced, for phosphide of copper, sulphide of lead, and other compounds still retain something of a metallic look. On the contrary, oxides and metallic chlorides (such as many compounds of phosphorus and sulphur) resemble salts.

All the other elements can be arranged in analogous series, more or less complete; for example, the silver series.

Atomic weights.

Ag=108; Cd=112; In=113; Sn=118; Sb=122;
Te=125 (?); I=127.

I shall only give the density of these bodies, because their concordance with the preceding series with regard to the other properties needs no explanation.

Density.

Ag=10.5; Cd=8.6; In=7.4; Sn=7.2; Sb=6.7;
Te=6.2; I=4.9.

(To be continued.)

ON THE SOLUBILITY OF SOLIDS IN GASES.*

By J. B. HANNAY, F.R.S.E., F.C.S., and JAMES HOGARTH.
(PRELIMINARY NOTICE.)

THIS investigation was undertaken in the hope that, by an examination of the conditions of liquid matter up to the "critical" point, sufficient knowledge might be gained to enable us to determine under what particular conditions liquids are dynamically comparable, in order that the microrheometrical method† (which the Royal Society has done one of us the honour of publishing in the *Philosophical Transactions*) might be applied to determine their molecular mass and energy relations. It seemed that as the laws relating to gases and liquids merge at what was called by Baron Cagniard de la Tour‡ "l'état particulier," and by Dr. Andrews§ the "critical point," an examination of matter up to the limit of the liquid state would be likely to yield us much information. The time we have to devote to scientific work being very limited, we found that it was quite impossible to make much advance by using the apparatus devised by Dr. Andrews, as the time required to change from one liquid to another was more than we had at our disposal. We therefore devised a new apparatus, which will be described in a more lengthy communication, but which, we may state, can be opened, the liquid changed, and again closed for a new experiment, in about one minute.

The question as to the state of matter immediately beyond the critical point being considered by Dr. Andrews to be at that time incapable of receiving an answer, we imagined that some insight might be gained into its condition by dissolving in the liquid some solid substance whose fusing point was much above the critical point of the liquid, and noticing whether, on the latter passing its critical point and assuming the gaseous condition, the solid was precipitated or remained in solution. We found that the solid was not deposited but remained in solution, or rather in diffusion, in the atmosphere of vapour, even when the temperature was raised 130° above the critical point, and the gas was considerably expanded. When the

side of a tube containing a strong gaseous solution of a solid is approached by a red-hot iron, the part next the source of heat becomes coated with a crystalline deposit, which slowly re-dissolves on allowing the local disturbance of temperature to disappear. Rarefaction seems to be the cause of this deposition, because if the temperature be raised equally and the volume retained at its original value, no deposition takes place. Those experiments have been done with such solvents as alcohol (ethyl and methyl), ether, carbon disulphide and tetrachloride, paraffins, and olefines, and such solids as sulphur, chlorides, bromides, and iodides of the metals, and organic substances such as chlorophyll and the aniline dyes. Some solutions show curious reactions at the critical point. Thus ethyl alcohol, or ether, deposits ferric chloride from solution just below the critical point, but re-dissolves it in the gas, when it has been raised 8° or 10° above that temperature.

It appeared to us to be of some importance to examine the spectroscopic appearances of solutions of solids when their liquid menstrua were passing to the gaseous state; but as all the substances we have yet been able to obtain in the two states give banded spectra with nebulous edges, we are only able to state that the substance does not show any appreciable change at the critical point of its solvent. Such was the case with anhydrous chloride of cobalt in absolute alcohol. It was suggested to us by Prof. Stokes that the substance obtained by the decomposition of the green colouring matter of leaves by acids, and which yields a very fine absorption-spectrum, might be useful for our purpose. We have prepared the substance according to the careful directions kindly furnished us by Prof. Stokes, and find that it shows the phenomenon in a marked manner, whether dissolved in alcohol or ether. The compound is easily decomposed by heat under ordinary circumstances, and yet can be dissolved in gaseous menstrua, and raised to a temperature of 350° without suffering any decomposition, showing the same absorption-spectrum at that elevated temperature as at 15° .

We considered that it would be most interesting to examine by this method a body such as sodium, which, besides being an element, yields in the gaseous state sharp absorption-lines. An opportunity seemed to be afforded by the blue solution of sodium in liquefied ammonia, described by Gore,* but we found that, on raising the ammonia above its critical point, the sodium combined with some constituent of the gas, forming a white solid, and yielding a permanent gas, probably hydrogen.

There seems, in some cases, to be a slight shifting of the absorption-bands towards the red, as the temperature rises, but we have as yet been able to make no accurate measurements.

When the solid is precipitated by suddenly reducing the pressure it is crystalline, and may be brought down as a "snow" in the gas, or on the glass as a "frost," but it is always easily re-dissolved by the gas on increasing the pressure. These phenomena are seen to the best advantage by a solution of potassic iodide in absolute alcohol.

We have, then, the phenomenon of a solid with no measurable gaseous pressure dissolving in a gas, and not being affected by the passage of its menstruum through the critical point to the liquid state, showing it to be a true case of gaseous solution of a solid.

Determination of the Elements of a Vibratory Movement: Measurement of Amplitudes.—M. Mercadier.—The author uses a so-called vibratory micrometer, a scale divided into millimetres, or fractions of a millimetre, intersected by a transverse line, extending to the zero of the scale, and forming with it a small angle. This may be either engraved upon the vibrating body itself, or upon a plate of glass, &c., fixed to the body so that this angular micrometer may share its vibrations, the scale being perpendicular to their direction.—*Comptes Rendus*.

* A Paper read before the Royal Society, November 20, 1879.

† On the Microrheometer, *Phil. Trans. Roy. Soc.*, 1879.

‡ *Ann. Chim.*, Series 2me, xxi., p. 127; xvii., p. 410.

§ "Bakerian Lecture," *Phil. Trans. Roy. Soc.*, 1869, p. 588.

* *Proc. Roy. Soc.*, vol. xxi., p. 145.

NON-PRODUCTION OF OZONE IN THE CRYSTALLISATION OF IODIC ACID.

By ALBERT R. LEEDS, Ph.D.

It has been stated by Prof. H. H. Croft* that air over crystallising iodic acid becomes ozonised. In preparing the acid according to Millon's process, Prof. Croft noticed that when the acid was left to crystallise over oil of vitriol the air in the jar (over the drying dish) had the characteristic odour and gave the usual tests of ozone. He states that the solution in every instance was boiled down to thin syrup, so that no trace of chlorine or nitric acid could possibly have remained. The air in the jar was tested from day to day, both by the smell and the action upon iodo-starch paper. Even when a few crystals began to form no change was noticed; but when the crystallisation set in fully the evolution was most remarkable, the strong smell being quite characteristic, and entirely different from that of chlorine or nitric acid.

We have repeated these experiments, and have found that the phenomena may be quite differently interpreted. After preparing the potassium iodate, by oxidation of iodine by means of potassium chlorate and nitric acid, and expulsion as far as possible of free chlorine from the solution, the potassium was converted into the barium salt, and the latter washed thoroughly. This was then decomposed by an equivalent weight of sulphuric acid, and, after separation from the insoluble salt, the solution of iodic acid was evaporated nearly to the point of crystallisation. It was then transferred to a retort, which was connected with three Geissler washing-apparatus, the first empty, the second and third containing water, and the last attached to a tube in which were placed ozonoscopes. Iodo-starch papers were likewise inserted in tubes placed in the neck of the retort. On aspirating a stream of desiccated air through the apparatus, the test-papers in the space above the iodic acid turned blue even before crystallisation set in. At the end of six hours they began to bleach, and at the expiration of twenty-four hours had become colourless. The test-papers exposed to the current after its washing through the small amount of water contained in the Geissler bulbs remained colourless throughout this and all succeeding trials. The strong odour given off from the crystallising solution was compared from time to time with the ozone generated by an electrical ozoniser, and was found to be essentially different in character. The crystallised iodic acid was then re-dissolved in the smallest possible amount of water, and re-crystallised in the same manner. The ozonoscopes in the space above the crystallising iodic acid were not affected on this repetition of the crystallisation. Now, when the difficulty of getting rid of every trace of extraneous matter by chemical operations—however carefully conducted—is borne in mind, the simplest explanation of the above phenomena, it appears to me, is that the apparent ozonic reaction is not due to ozone produced in the act of crystallising,—which, as Prof. Croft remarks, is anomalous—but to a trace of chlorine or nitrous acid, or possibly some lower oxide of iodine formed in the process of manufacture, and eliminated by successive crystallisations of the acid. The fact that the air after washing did not manifest the ozone reaction strongly corroborates this view.

NEW AND OLD VIEWS ON THE NASCENT STATE OF BODIES.

By Dr. T. L. PHIPSON.

IN my note on this subject, in the CHEMICAL NEWS (vol. xl., p. 184), I called attention to some statements by Prof. Tommasi published in this journal (vol. xl., p. 171). The

* CHEMICAL NEWS, vol. xxv., p. 87; American Journal of Science, III., p. 466; Canadian Journal, xiii., No. 3.

author has since replied in an interesting paper (vol. xl., p. 245), and it is evident that our opinions on the nascent state of bodies are not quite so divergent as his first note led me to expect.

Many chemists will doubtless have been misled by the title of Dr. Tommasi's first note, "On the Non-Existence of Nascent Hydrogen"; but it appears by his second note that he admits its existence, for he says:—"The reductive power of nascent hydrogen varies according to the chemical reaction which produces it. And if this gas in the nascent state possesses greater affinity than in the natural state, it is solely due to the fact that the hydrogen, the moment it issues from a combination, is found to be accompanied by the whole quantity of heat produced during the setting free of the hydrogen." So that the author not only does not deny the existence of nascent hydrogen, but actually supports it by a theory of his own. This is all I contended for.

Dr. Tommasi hopes that I may accept his thermic theory. It is, perhaps, as good as another; but if the word *heat*, in the paragraph just quoted, be replaced by *electricity*, this thermic theory becomes identical with that published in 1858 in my "Force Catalytique, &c." (quoted in my previous note). Whether the thermic theory will explain such facts as those given on pages 29, 30, and 31 of my paper better than the electric theory still remains to be proved; whilst the paragraph on page 34 of that work will convince him that I was occupied with this subject as early as 1857; that is, not only before Tommasi, but before Houzeau.

But by whatever theory these curious reactions may be explained, there can be no doubt that Dr. Tommasi has made known several new and very interesting facts concerning reductions operated by hydrogen; and he will be the last person not to see the identity of the reduction of auric chloride by hydrogen gas in presence of platinum, for instance, with the phenomena of hydrogen in the nascent state.

His argument that *heat* will produce the same result, when the temperature of the hydrogen gas is raised, is rather old,—it will also be found in my "Force Catalytique" already quoted—but we may use the same argument and substitute the word "*electricity*."

After all what is *caloric*? It is not a material substance that can be introduced into a body like raisins into a cake. The electric action can be put in evidence by the galvanometer more directly than the thermic condition can be rendered manifest by the calorimeter; and, as caloric and electricity are mutually convertible agencies, replacing each other equivalent for equivalent, as Grove and others have long since shown, it is no doubt difficult to admit that either one or the other is, in all cases, essentially the cause of the "nascent state" of bodies.

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ON CLARKE'S METHOD FOR THE SEPARATION OF TIN FROM ARSENIC AND ANTIMONY.

By FRED. P. DEWEY.

IN the beginning of the year 1876 an investigation was undertaken on the separation of tin, arsenic, and antimony. After some study on the published methods, and some work on a few of them, it was decided to make a thorough investigation of the method proposed by F. W. Clarke,* as that seemed the most favourable method, notwithstanding the poor results obtained by G. C. Wittstein.† Clarke's method is to add to the solution of the metals prepared as for the precipitation of sulphides, oxalic acid in the proportion of twenty parts to one of tin, keeping the solution so concentrated that oxalic acid will separate in the cold.

* Am. Jour. Sci., 49, 145.

† Fresenius's Zeitschrift, 9, 487.

Hydrogen sulphide gas is then passed through the boiling solution until the antimony and arsenic are all precipitated. The solution is allowed to stand in a warm place for half an hour, the sulphides are filtered off, and separated according to any of the well-known methods. The filtrate is neutralised by ammonia, the precipitate formed dissolved in ammonium sulphide, and acetic acid added in excess, whereby the tin is precipitated as a light coloured mixture of sulphide and oxide, which soon darkens in colour. When complete precipitation is ensured by standing in a warm place several hours, it is filtered off, washed in ammonium nitrate solution, ignited, and weighed as stannic oxide, SnO_2 . The results given by Clarke were the recovery of 99.93 and 99.57 per cent of the tin from solutions containing unknown amounts of antimony and arsenic, the latter result being considered poor by him, and owing to incomplete precipitation from filtering too soon after the addition of reagents. It was originally intended to investigate the separation of tin from antimony and tin from arsenic separately, and then tin from antimony and arsenic; but it soon became evident that, with the time at my disposal, I could only undertake the separation of tin from antimony, and the experiments here described were devoted entirely to that object.

I made two separations, intending to follow Clarke's plan exactly. Pure metallic antimony was weighed out and dissolved in strong hydrochloric acid with the addition of potassium chlorate. As soon as all the antimony had gone into solution, weighed amounts of pure metallic tin were added, and the final solution evaporated to dryness to drive off hydrochloric acid and decompose all potassium chlorate: the residue was taken up in a few drops of hydrochloric acid and water, and the requisite amount of oxalic acid added. As oxalic acid is soluble in 10 parts of cold water,* one solution could not be over 55 c.c. and the other over 39 c.c., no account being taken of the few drops of hydrochloric acid added. Hydrogen sulphide was transmitted through the boiling solution for half an hour, when the antimony seemed to be all down. After standing for half an hour in a warm place, a portion of the fluid was run through a weighed filter, and the filtrate tested with hydrogen sulphide, when more antimony came down; this was filtered off, and the second filtrate tested, when still more antimony came down. Thereupon I gave up the idea of keeping the solution so concentrated, and diluted it a little, when the antimony came down, so that on testing the filtrate by diluting and adding more hydrogen sulphide no antimony was precipitated. The precipitate was filtered off, washed well, dried at 100°C ., and weighed. The filtrate was diluted largely, ammonia added, the slight precipitate dissolved in ammonium sulphide, acetic acid added in excess, and the whole allowed to stand over-night, when the precipitate was filtered off, washed in ammonium nitrate, ignited, and weighed as stannic oxide, SnO_2 :—

Used.		Found.	
Sb.	Sn.	Sb.	Sn.
0.3089 gr.	0.2723 gr.	0.3063 gr.	0.2704 gr.
0.2671 "	0.1908 "	0.2704 "	0.1821 "

In a second set a portion of the antimony trisulphide was heated to 230°C . in a tube filled with dry carbon dioxide, and the loss calculated for the whole precipitate:—

Used.		Found.	
Sb.	Sn.	Sb.	Sn.
0.1542 gr.	0.2705 gr.	0.1356 gr.	0.2717 gr.
0.1172 "	0.3736 "	0.1186 "	0.3760 "

Some qualitative experiments were undertaken to test the influence of free hydrochloric acid on the separation, and also the statement of Clarke that antimony could not be detected in the filtrate from the antimony trisulphide

either by the Marsh test or the black stain on platinum with zinc, and that oxalic acid did not interfere with either of these tests. I found that oxalic acid obscured the platinum and zinc test by the formation of a dense white coating on both the platinum and zinc when the acid became nearly neutralised (probably an oxalate of zinc), which completely masked any black stain that might have been produced on the platinum, but that it did not obscure the Marsh test in the least. Five solutions were prepared as before, each solution containing 0.2 gr. tin and 0.15 gr. antimony, and evaporated to dryness, the residues being taken up in oxalic acid; and, starting with no free hydrochloric acid, 1, 3, 5, and 7 c.c. of hydrochloric acid of 1.09 sp. gr. were respectively added. These solutions were diluted to 250 c.c. Hydrogen sulphide was transmitted through the boiling solutions for three-quarters of an hour, and the antimony trisulphide filtered off. The filtrates were tested in an ordinary Marsh apparatus.

No. 1 containing no HCl gave only a slight yellowish mirror, probably S.

No. 2 containing 1 c.c. HCl gave a metallic mirror for about two minutes.

No. 3 containing 3 c.c. HCl gave a good metallic mirror for some time.

No. 4 containing 5 c.c. HCl gave a very good metallic mirror for a long time, about equal portions of the filtrates being used for each test.

No. 5 containing 7 c.c. HCl deposited thin red flakes of Sb_2S_3 on concentration.

These results show that to get the best separation no hydrochloric acid can be present, although a very small amount can be present without exerting any very great solvent action on the antimony trisulphide. The precipitates of antimony trisulphide were dissolved in strong hydrochloric acid, evaporated, and treated with zinc: the metallic precipitates were treated with small amounts of dilute hydrochloric acid, filtered, and the filtrates tested with mercuric chloride, when heavy precipitates of mercurous chlorides came down in every case. These results show that it is always necessary to make double precipitations of the antimony, a point suggested by Clarke as being necessary only when great accuracy is required. The results of the above described experiments indicate that the following modifications of the original plan are required, viz., a solution *without free mineral acids*, and diluted to 250 c.c. for 0.2 gr. of each metal; also, in all cases a double precipitation of the antimony.

Solution in hydrochloric acid with potassium chlorate being rather tedious, a mixture recommended by Dr. Clemens Winckler, *Fresenius's Zeitschrift*, consisting of 1 part strong nitric acid, 4 parts strong hydrochloric acid, and 5 parts water, was tried and found very satisfactory. The evaporation of a solution of tin and antimony prepared by this mixture cannot safely be undertaken, on account of the well-known fact that stannic chloride will volatilise from a dilute acid solution, and both stannic chloride and antimony trichloride from a concentrated acid solution on evaporation, and the use of an alkali to neutralise the acid, is objectionable for reasons I shall state beyond; but from the fact that in the analyses given the metals were dissolved in hydrochloric acid and potassium chlorate and evaporated to dryness without any very great loss on the total amount of metals used, it suggested itself that the presence of a sufficient amount of potassium chloride to form double salts with all of the tin and antimony used would serve to fix the tin and antimony during evaporation. Two experiments were made to ascertain definitely whether tin or antimony would volatilise under such conditions. 1.7 grs. of tin were dissolved in 50 c.c. of the aqua regia and 2.5 grs. of potassium chloride were added. The solution was transferred to a retort of 350 c.c. capacity, diluted to 100 c.c., and distilled at about 100°C . in an air-bath, while a slow current of air was forced through the retort. When half of the fluid had passed

* Storer's "Dictionary of Solubilities."

* The poor results obtained by Wittstein were probably due to the presence of large amounts of free acid in his solutions.

over, the receiver was removed, emptied, and returned, then the distillation was resumed. The distillate was diluted to 100 c.c., part of the acid was neutralised by sodium carbonate, and hydrogen sulphide was transmitted through it, when there was only a slight separation of sulphur, probably caused by chlorine given off from the aqua regia. The distillation was carried to dryness, the distillate treated in the same way, and with the same result. A second portion of tin was treated in the same way, omitting the addition of potassium chloride, and in each portion of the distillate heavy precipitates of tin sulphide were obtained. 1.5 grs. of antimony were treated exactly as the tin in the first experiment with the same result. 1.5 grs. more of antimony were treated without potassium chloride. In the first distillate, containing three-quarters of the liquid, no antimony could be detected, but in the final distillate a large amount of antimony was found. These experiments show that solutions of stannic chloride and antimony trichloride containing free hydrochloric or nitric acid, or both, can be safely evaporated to dryness if a sufficient amount of potassium chloride is present. In all of my subsequent analyses the free acid was removed by evaporation to dryness in the presence of an excess of potassium chloride.

The course next adopted was to dissolve about 0.2 gr. of each metal in the aqua regia, add sufficient potassium chloride, and evaporate to dryness. The requisite amount of oxalic acid was then placed in the beaker, and boiling water added until a clear solution was obtained, which is very easy to do. The solution was diluted to 250 c.c. Hydrogen sulphide was transmitted through the boiling solution for half an hour. The precipitate was immediately filtered off* and washed. The precipitate of antimony trisulphide containing a little tin sulphide was dissolved in ammonium sulphide. This solution was poured into a boiling solution of oxalic acid, containing enough of the acid to decompose the ammonium sulphide, and still have sufficient excess to hold the tin in solution. Hydrogen sulphide was transmitted through the boiling solution for ten minutes, and the solution filtered through a weighed filter. The precipitate was well washed with hot water, dried at 100° C., weighed, and a portion converted into pure anhydrous antimony trisulphide as before. The filtrates containing the tin were united and treated with the proper reagents. The precipitate was filtered off after standing, washed in ammonium nitrate, and weighed as tin oxide. Each filtrate was always tested to ensure complete precipitation:—

Used.		Found.	
Sb.	Sn.	Sb.	Sn.
0.2243 gr.	0.2238 gr.	0.2390 gr.	0.2218 gr.
0.1908 "	0.2346 "	0.2058 "	0.2219 "
0.3871 "	0.2573 "	0.3949 "	0.2607 "
0.3108 "	0.2187 "	0.3158 "	0.2203 "

Repeated testings of the weighed tin and antimony indicated that the actual separations were satisfactory. There were, however, some difficulties and some obvious sources of error in the methods used for their determinations which would cause gain of antimony and gain or loss of tin. On dissolving the antimony sulphide in strong hydrochloric acid a black residue was always obtained, no matter how much the precipitate had been washed, indicating the presence of oxalic acid in the precipitate in some form. The solution of tin required extreme dilution to get it all down, the solution being diluted in one case to nearly 1500 c.c., which seemed to be due to the unavoidable presence of large amounts of ammonium oxalate. Besides this, the washing of the tin precipitate was very

* It was observed that when the fluid with the Sb_2S_3 precipitate was allowed to stand to remove H_2S , there was always a separation all through the liquid of flakes which did not look like S, but did look like SnS_2 ; a supposition supported by the fact that these flakes separated in the filtrate from Sb_2S_3 , filtered off immediately as it cooled during filtering, and were dissolved again on applying heat.

difficult and tedious indeed, which is a reason for keeping the amount of salt in the solution as small as possible.

To avoid the difficulty of precipitating the tin completely and washing it thoroughly, it seemed necessary to remove the oxalic acid entirely from the filtrates containing it. Strong sulphuric acid was decided upon as the best means—that is, to evaporate the solution to dryness, and treat with strong sulphuric acid at about 100° C., which rapidly decomposes the oxalic acid into water, carbon monoxide and dioxide. In two experiments 0.5 gr. tin was dissolved in the aqua regia, potassium chloride was added, the solutions were evaporated to dryness, and the residues taken up in oxalic acid, 25 grs. in the one and 30 grs. in the other. They were again evaporated to dryness, and treated with about 40 c.c. of strong sulphuric acid at 100° C. until all effervescence had ceased, when they were diluted to 500 c.c., treated with hydrogen sulphide gas for half an hour and allowed to settle, after which the precipitates were filtered off, washed in ammonium nitrate, and weighed as tin oxide, yielding 0.4998 and 0.4996 gr. tin. Applying this process to the filtrate from the antimony trisulphide, the following separations were next made, the process being conducted as before except the dilution of the solution before the precipitation of the antimony trisulphide to 400 c.c., when 0.5 gr. of each metal were used:—

Used.		Found.	
Sb.	Sn.	Sb.	Sn.
0.2000 gr.	0.2000 gr.	0.2084 gr.	0.1964 gr.
0.2000 "	0.2000 "	0.2030 "	0.2031 "
0.3000 "	0.3000 "	0.3070 "	0.3029 "
0.5000 "	0.5000 "	0.5066 "	0.5004 "
0.5000 "	0.5000 "	0.5177 "	0.4962 "

The last two antimony results were corrected for the carbonaceous residue, which was filtered off on a small filter and weighed, being 0.0085 and 0.0070 gr. Following the analogy of the two cases, we may suppose, as J. P. Cooke, jun.,* does in his determination of the atomic weight of antimony, that this residue comes from occluded oxalic acid.

A slight modification of Bunsen's† method of weighing antimony as antimony tetroxide having given 0.4996 and 0.4986 gr. antimony instead of 0.5000 gr., four separations were made in which the antimony was weighed as antimony tetroxide, the other details remaining as before:—

Used.		Found.	
Sb.	Sn.	Sb.	Sn.
0.5000 gr.	0.5000 gr.	0.5190 gr.	0.4911 gr.
0.5000 "	0.5000 "	0.5067 "	0.4963 "
0.5000 "	0.5000 "	0.5136 "	0.4973 "
0.5000 "	0.5000 "	0.5146 "	0.4926 "

While the results obtained by this method are no better than those obtained by weighing antimony as trisulphide, yet it required less time, and is a much neater operation to get them.

Since reaching the above results and conclusion Bunsen‡ has published further experiments on this method of weighing antimony, and reaches the conclusion that it cannot easily be done with accuracy, because more oxygen than is necessary will be given off at a temperature only slightly above that required to convert antimony pentoxide into the tetroxide, and at a very intense heat the trioxide thus formed will be volatilised. From my results it would seem that the heat employed was too low rather than too high.

In conclusion, I desire to express my thanks to Prof. O. D. Allen for much kind assistance during the progress of the above work.—*American Chemical Journal*.

* *Transactions of Amer. Acad. Sci., Boston*, 13, 15.

† *Ann. der Chemie*, 106, 3.

‡ *Fresenius's Zeitschrift*, 18, 268.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, November 20, 1879.

Dr. GILBERT, Vice-President, in the Chair.

THE minutes of the preceding meeting were read and confirmed.

THE CHAIRMAN announced that a ballot for the election of Fellows would take place at the next meeting of the Society (December 4).

THE following certificates were read for the first time:—
J. Howard, J. Knowles, J. Snodgrass, A. Scott, E. J. Day, J. Steiner.

THE CHAIRMAN then called on Mr. CHURCH to read a paper. "*A Chemical Study of Vegetable Albinism (Part II., Respiration and Transpiration of Albino Foliage)*." The author has proved that white foliage does not possess the power, even in sunshine, of decomposing the carbon dioxide of the air, but adds largely to the normal amount of that gas in the air, thus resembling the petals of flowers and the action of green leaves during darkness. The best results were obtained with the maple (*Acer negundo*), the holly, the ivy, and the *Alocasia macrorrhiza*. Every care was taken to choose white and green foliage comparable in every respect, age, &c. The leaf stalks were placed in boiled distilled water, and covered by a bell jar inverted over mercury. To the mouth of this bell jar is fitted a cork, through which pass two tubes, one long and the other short. Air is sucked from the long tube through a Pettenkofer tube containing baryta water. At the end of the experiment the Pettenkofer tube is washed out without exposure to the atmosphere, and the carbonate determined after the method of Dupré and Hake (*Chem. Soc. Journ.*, March, 1879). The CO₂ in the atmosphere was found to be 3.21 parts in 10,000. The author embodies his results in a table. 1000 sq. centimetres of the white foliage of the maple evolved, in two hours, 16.69 parts of CO₂ per 10,000 of air; 1000 sq. centimetres of green foliage evolved 0.44 parts CO₂. Similarly, 1000 sq. centimetres of white holly leaves evolved 18.82 parts CO₂ per 10,000; of green, 4.49. 1000 sq. centimetres of white *Alocasia* leaves in two experiments evolved 15.06 and 38.96 parts of CO₂ per 10,000; of green, 1.14 parts. The author has also studied the comparative loss and gain of albino and green foliage. White holly sprays placed in water gained, in two hours, 0.29 per cent; green holly, under similar conditions, gains 1.55 per cent. When no water was supplied the white holly lost 0.54 per cent; the green, 10.26 per cent. The author promises further work on this interesting subject. He exhibited some dried specimens of albino and ordinary leaves, the albino leaves being thinner and altogether much less substantial in their structure.

DR. GILBERT said that the results obtained by the author seemed to be of considerable importance. In a former paper the composition of the green and white leaves had been investigated, the white leaves containing less lime but more non-albuminoid nitrogen than the green, indicating a capability for development without actual growth. In the present paper the author shows that these white leaves do not perform leaf functions.

MR. WARINGTON remarked that in the white leaves respiration seemed to take place in an exalted state. He would like to know whether much starch had been found in the white leaves?

MR. GROVES asked if there was any difference in the number of stomata in the white and green leaves?

DR. VOELCKER enquired where the large amount of carbonic acid given off by the white leaves came from? The researches seemed to indicate that the white leaves seemed to be partially dead; if not quite deceased, at least

gradually decomposing. This would account for the evolution of CO₂.

MR. CHURCH, in reply, said there was more starch in the green than in the white leaves; that the leaves used in the experiments were all in active growth and not in the least decaying. In his opinion the CO₂ given off by the white leaves came from substances elaborated by the green leaves. In the same way the flowers of the white lily and green leaves in the dark evolved CO₂ without any actual decay. The green leaves gave off CO₂ as well as the white, but in sunshine they decomposed more than they excreted. As far as he knew white and green leaves had the same number of stomata.

MR. KINGZETT then read a paper entitled, "*Contributions to the History of Putrefaction*," Part I. The author discusses to a certain extent the process of putrefaction, and concludes that it would naturally be expected that a substance allowed to undergo putrid fermentation without oxidation would more readily undergo chemical oxidation than the original integral mass, and therefore that the oxygen absorbing capacity of a substance would progress increasingly with such putrefactive decomposition. He therefore endeavoured to determine the oxygen-absorbing capacities of organic solutions from time to time as they passed into putrefaction, and to compare these numbers with the oxygen capacities of similar solutions protected during the same periods by so-called antiseptics. Various difficulties were met with, and the author does not offer all the results obtained. Some of the experiments, however, have an important bearing on Tidy's oxygen process, which, in the words of its author, "undoubtedly furnishes us with exact information as to the relative quantities of putrescent and easily oxidisable matter, and of non-putrescent and less easily oxidisable matters present in the water." The author of the present paper thinks that the oxygen process is liable to mislead chemists in interpreting their results if the most pernicious organic matters are those which are in a putrescent state; for, from the results obtained by Kingzett, it appears that the oxygen absorbed by a water containing organic matter in a non-putrescent condition is more than that required by the same organic matter in a putrescent state. Thus, 5 c.c. of a dilute solution of albumin, which began to stink after 150 hours, required, when fresh, permanganate equal to 0.008004 grm. oxygen; after keeping 24 hours, 0.008671; after 120 hours, 0.0076038; after 1104 hours, 0.00667; after 1176 hours, 0.0062031; after 1440 hours, 0.0058696; after 1534 hours, 0.0054694; showing, after a slight rise (8 per cent), a decrease of over 30 per cent in the oxygen absorbed. Similar results were obtained with extracts of beef, fish, &c. Thus the author concludes that the oxygen process might pass a water containing pernicious products, since it is possible to obtain an organic solution in an active state of putrefaction, which requires less permanganate than the same solution in a fresh state.

PROF. TIDY said that the results obtained by Mr. Kingzett required thinking about. No doubt different materials had different oxygen-absorbing powers. As a practical man he was satisfied with the third place of decimals. He was pleased to see Mr. Kingzett found it necessary to go to the seventh.

MR. OTTO HEHNER thought that the results rather proved than disproved the accuracy of the oxygen process, because the oxygen-absorbing power did at first increase; afterwards, of course, substances nitrified, and in other ways the organic matter disappeared. This rise and subsequent decrease he had himself pointed out, using the albuminoid ammonia process.

MR. CHURCH suggested that some inaccuracy might arise from the deposit which fell as the solution putrefied, the successive quantities of 5 c.c. containing less and less organic matter as the deposit increased.

AFTER some remarks by Dr. VOELCKER and a brief reply by Mr. KINGZETT to the various points raised in the discussion,

THE next paper was read by Dr. WRIGHT. It was en-

titled, "Notes on Manganese Dioxide," by C. R. A. WRIGHT and A. E. MENKE. The authors have repeated many of the experiments of former observers in some cases very different results. They have made many careful gravimetric analyses of the substances precipitated under various conditions, the experiments being more particularly directed to the examination of substances formed in presence of potassium compounds so as to determine how far potash is carried down in combination with the manganese. Substances were prepared by the action of nitric acid or sulphuric acid on potassium permanganate; by solution in hydrochloric acid and reprecipitation by water in presence or absence of potassium salts; by acting on potassium permanganate with sulphurous acid, alcohol, and glycerin; by precipitating manganese salts with potassium permanganate under various conditions; by acting with air and caustic potash on manganese salts (Weldonising); by precipitating from solution by bromine, &c., under various conditions; and by heating manganese nitrate to 160° alone and in the presence of potassium nitrate. In every case except the last potassium was present in the ultimate product in considerable quantity, even though the precipitate was formed in presence of very large quantities of free hydrochloric, sulphuric, nitric, or acetic acid. Other metals if present are often carried down in preference to potassium, e.g., calcium. Unless the circumstances of the experiment are such as to allow of the precipitation of a large percentage of potassium or other equivalent metal, the precipitate never contains as much oxygen as corresponds to MnO_2 , the deficiency being greater or less as the quantity of potash present is less or more. The only exception to this rule occurs when hot solution of manganese salt and excess of permanganate act on one another; with cold solutions the precipitate is deficient in oxygen unless zinc sulphate be added. Manganese dioxide containing potash can be completely converted into MnO , retaining but traces of potash by long-continued ignition in a current of hydrogen. The green potash-free oxide thus formed does not spontaneously oxidise in the air. When freshly thrown down and just dry to the sight and touch, the dioxide is a hydrate, $4MnO_2 \cdot H_2O$; on standing over sulphuric acid this hydrate loses water indefinitely and continuously. $MnO_2 \cdot H_2O$ is the lowest hydrate stable in dry air. Up to 108° no appreciable loss of oxygen accompanies the expulsion of water. At 210° all the precipitated oxides, &c., tried lose oxygen slowly and regularly. The nearly anhydrous MnO_2 , prepared by heating manganese nitrate to 160°, does not evolve oxygen at 210°. In all cases perceptible amounts of water from a few tenths to two per cent are retained after many hours drying in a current of dry pure air at 210°. The volumetric processes of Kessler and Pattinson have been examined; they give good results. The authors suggest a convenient modification of Pattinson's method. The diluted solution, slightly acid, is heated to nearly 100°, a considerable excess of bromine water added, and then freshly precipitated zinc carbonate, all free bromine being expelled by boiling; ferric chloride usually forms permanganate, and thus complicates the manipulation. A more convenient and quite as accurate a method as any of those requiring bromine, &c., was found to be the addition of the diluted and not too acid solution of manganese in the cold to a small excess of dilute permanganate solution containing zinc sulphate. The precipitate is collected on a glass-wool funnel, dissolved in excess of acid ferrous sulphate, and titrated as usual. Three-fifths of the manganese in the precipitate represent the manganese in the original solution. The authors have verified the statement of Gorgeu that actual MnO_2 is obtained by heating manganese nitrate to 160°. The statements of Guyard as to the effect of heat on the precipitate formed by adding manganese salts to permanganate are also corroborated. The recent statements of Pickering as to the deficiency of oxygen in the substances thrown down by acting with water on the superchlorides of manganese are also substantially correct. The accuracy

of the results of Volhard as to obtaining pure MnO_2 free from potash by precipitating manganese sulphate with permanganate in the presence of nitric acid, &c., is questioned by the authors. The experiments of Morawski and Stiegl have not been confirmed as far as they have been repeated, viz., hydrated manganese dioxide containing potash does not (as stated by these chemists) lose all its water by ignition. The substance thrown down by the action of manganese sulphate on potassium permanganate is not $3MnO_2 \cdot 2H_2O$, but an indefinite hydrate containing much potash. On boiling with caustic potash it does not form (after washing and drying at 100°) $Mn_4KH_3O_{10}$ or $8MnO_2 \cdot K_2O \cdot 3H_2O$ as stated, but a substance much richer in potash, $5MnO_2 \cdot K_2O$. $Mn_4KH_3O_{10}$ does not uniformly result by the action of alcohol, glycerin, &c., on potassium permanganate, but, instead, substances of very variable composition. The authors also notice incidentally that manganese chloride is sensibly volatile at a red heat in a current of HCl, and that perceptible errors in alkali determinations may be introduced if solutions containing ammonium sulphide are evaporated down in glass or porcelain. Identical results were obtained when titrating the available oxygen in a substance, whether the material was dissolved in acid ferrous sulphate checked by dichromate, or was distilled with HCl into KI, and the titrated iodine estimated by adding excess of thiosulphate and titrating back with iodine solution.

The next paper was read by the SECRETARY, "On the Reaction between Sodium Thiosulphate and Iodine: Estimation of Manganese Oxides and Potassium Bichromate," by S. PICKERING. In a former paper the author suggested a modification of Bunsen's volumetric method, viz., transferring the sample to be analysed to a beaker containing a large excess of potassium iodide, adding a small quantity of acid, and determining the iodine liberated by running in sodium thiosulphate solution. This method gives results slightly higher than the original method of Bunsen, viz., boiling the oxides with hydrochloric acid, and collecting the evolved chlorine in a solution of potassium iodide. The author has in the present paper exhaustively examined the effects produced by varying the several conditions of the reaction. When iodine and sodium thiosulphate react in warm solution a decomposition,—



attended with the formation of sulphate occurs, as well as the ordinary reaction, $I_2 + 2Na_2S_2O_3 = Na_2S_4O_6 + 2NaI$. Thus at 52° 3.9 per cent of iodine reacted to form sulphate; at 0° 1.84 to 0 per cent reacted in a similar manner. Dilution increases very slightly the amount of sulphate formed. An excess of potassium iodide is without effect. Dilute solutions of iodine do not vary in four days as regards the amount of iodine they contain, whether kept in the dark or in diffused daylight. Within certain limits the amount of iodine liberated at once is proportional to the amount of hydrochloric acid added, but the acid has no influence on the relative proportions in which the two above-mentioned reactions take place. No difference occurs whether the reaction is made by adding iodine solution to an excess of thiosulphate or thiosulphate to the iodine; if, however, hydrochloric acid be present the amount of sulphate is increased. The author discusses the various forms of apparatus for estimating the oxides of manganese by Bunsen's plan: he prefers boiling the oxide in a small flask fitted with a thistle funnel, the chlorine being absorbed in three other flasks containing potassium iodide. Traces of chlorine were retained by the acid liquid, even after prolonged boiling. If undiluted acid be used the Bunsen method gives results almost identical with those obtained by using the plan suggested by the author; if the acid be diluted the amount of chlorine evolved is diminished in the Bunsen method. If potassium bichromate be estimated by the Bunsen method a notable deficiency of chlorine is observed, even if the acid be undiluted. If chlorine-water be diluted and boiled,

considerable loss of chlorine takes place from the formation of hydrochloric acid. This loss begins at low temperatures. This probably is sufficient to explain the low results obtained by the Bunsen method. The author's modification is, of course, not applicable to manganese ores owing to the presence of ferric oxide. Bunsen's method must therefore be used with the precautions above indicated.

The Society then adjourned to December 4, when a ballot for the election of Fellows will take place, and the following papers will be read:—"On the Theory of Fractional Distillation, Part II.," by F. D. Brown; "On the Influences Exerted upon certain Chemical Changes by Variations in the Amount of Water of Dilution," by M. M. P. Muir and C. Slater; "On α - and β -phenanthrene Carbonic Acid," by F. R. Japp; "On some Derivatives of Phenyl-acetic Acid," by P. Philippa Bedson.

PHYSICAL SOCIETY.

Ordinary Meeting, November 22, 1879.

Prof. W. G. ADAMS in the Chair.

New members:—Prof. Reilly and Prof. Heath, of Cooper's Hill Engineering College.

Prof. GUTHRIE exhibited a new Image Retention Photometer, and demonstrated its action to the meeting. It consists of two fixed plane mirrors inclined to each other at an angle of 45° . The rays from the two sources of light to be compared are allowed to fall on these mirrors, those from one source, say that on the right hand, falling on the right-hand mirror, and those from the left-hand source on the left-hand mirror. These rays are again reflected from the mirrors at right angles to their former paths, and thrown upon a semi-transparent screen, where their relative intensities can be compared by the eye of the observer. Between the mirrors and each source of light a revolving shutter is interposed. These shutters are formed of brass disks, and they are both mounted on the same axis, which can be turned by hand or otherwise. They would completely screen the light from the mirrors were it not that each is provided with four radial apertures or slots, through which the rays can pass. The slots on the side at which the brighter source of light is placed are narrower than those on which the weaker source is placed. The latter slots are made adjustable in size by sliding blinds, and a scale is added to measure the degree to which they are closed. On revolving the shutters the reflection of the rays to be compared are seen side by side and (owing to the persistence of images on the retina) continuously on the screen placed in front; and they are brought to equality of brightness by closing or opening the blinds of the adjustable shutter. When this is so the ratio of the respective orifices of the shutters as given by the scale is the inverse ratio of the luminous intensities compared.

Prof. ADAMS remarked that the speed of rotation should be such as to produce a uniform field of light on the screen, a result which hand-turning was not very conducive to.

Prof. FOSTER observed that the use of this new photometer might be less fatiguing to the eye than those photometers which presented a steady beam to the eye undiluted with intervals of darkness, during which the light is cut off, as in the instrument before the meeting.

Prof. REINOLD then read a paper by Prof. RÜCKER, of the Yorkshire College, Leeds, "On a Suggestion as to the Constitution of Chlorine offered by the Dynamical Theory of Gases." If a gas of density, δ , consists of molecules, each of which possesses m degrees of freedom, and if also the inter-molecular forces are negligible, the specific heats at constant pressure (C_p) and at constant volume (C_v) are connected by the two well-known equations—

$$(1.) \quad (C_p - C_v) \delta = 0.694.$$

$$(2.) \quad \frac{C_p}{C_v} = 1 + \frac{2}{m+e}.$$

Where e is a quantity which depends upon the potential energy of a molecule. Hence, if C_p is given by experiment, C_v can be calculated from the first equation, and then $m+e$ is known from the second. Regnault determined the specific heats at constant pressures of 35 gases, and from the experiments of E. Weidemann and of Wullner it appears that his values are correct within 6 per cent, and that $m+e$ can be calculated very approximately from the above equations if C_p is given. One of the chief difficulties of the thermo-dynamic theory of gases has been to attribute to m and e values which would at once lead to the observed ratios of C_p and C_v , and satisfy any rational supposition as to the interior mechanism of a molecule. Kundt and Warhng proved that for mercury—

$$\frac{C_p}{C_v} = 1.666,$$

which is consistent only with the supposition that the atoms of mercury are smooth rigid spheres; and Boltzmann and Bosanquet have pointed out that for a smooth rigid surface of revolution—

$$\frac{C_p}{C_v} = 1.4,$$

a number agreeing closely with the experimental values for air, O, N, H, CO, and NO. The molecules of these gases may therefore be constituted of two spheres rigidly united, or, as Mr. Rücker suggests, bound together by forces which prevent the separation of their surfaces while leaving them otherwise free to move. The principal object of Prof. Rücker's paper was to point out an interesting fact connected with the application of this theory to chlorine. The maximum number of degrees of freedom which a molecule composed of smooth & rigid spheres could possess would be $3n$, but the forces in play between the spheres might reduce the number. Thus, the value of $m+e$ could not exceed, but might be less than, $3n+e$. When the molecule consists of two atoms $e=0$, but for more complex molecules we should, *ceteris paribus*, expect its value to increase with the number of molecules. From two tables of results calculated by him, Prof. Rücker, however, finds that for a number of simple and complex gases and vapours the value of $m+e$ is for each substance less than $3n$ (or the maximum possible value of m), while for the majority of the chlorine compounds examined the reversed statement holds good; that is, the value of $m+e$ is generally greater than $3n$. This difference may be explained by supposing that for chlorine e is abnormally large, and that the spheres are not necessarily in contact; or that n has been taken too small, that the symbol Cl_2 is incorrect, and that the chlorine atom contains a larger number of sub-atoms than has been supposed, a supposition which accords with the recent researches of Prof. Victor Meyer on the vapour-density of chlorine. Prof. Rücker also finds that the ratio of the specific heats of bromine and one of its compounds studied ($\text{C}_2\text{H}_2\text{Br}$) agrees with those of chlorine and the corresponding chlorine compound.

Dr. SHETTLE, of Reading, then read a paper on the "Influence of Heat upon certain forms of Induction Coils, considered more especially in relation to the Inductive Power which the Blood Exercises on the Various Structures of the Body." The author found that when a copper and a zinc wire were insulated from each other by parchment-paper and paraffined silk, and wound in close proximity to each other, an (induced) current was indicated on a galvanometer whose terminals were connected to the neighbouring ends of the zinc and copper wires respectively, the other ends being left free. When the latter were connected across the deflection was *nil*. On raising the temperature of the two wires by causing hot water to flow inside the coil into which they were wound the deflection was largely increased. These experiments led Dr. Shettle to imagine that there is a similar action in the animal body. The heart is made up of nerves and muscular fibres winding spirally, and some of these wind round each other as as

to form a spiral cord, round which the blood capillaries also wind. Dr. Shettle compares these nerve and muscle bundles to the coils of zinc and copper wire in his experiments, and infers that electric currents may be induced in them as in the wires. The flow of the warm magnetic blood would also tend to produce currents in them. Dr. Shettle, further, drew attention to the fact that animals live and move in a magnetic field, and that electricity must be generated in them by their movements, internal and external.

Mr. ENMOTT exhibited Crossley's Form of Microphone, which consists of four short rods of carbon jointed loosely into four blocks of carbon, so as to form a square. It is used as a transmitter for telephones, and Mr. Crossley regularly transmits the services of a church with it to several hearers. Its speaking, singing, and whistling powers were successfully demonstrated to the meeting.

CORRESPONDENCE.

CARBON IN STEEL.

To the Editor of the Chemical News.

SIR,—M. Sergius Kern (CHEMICAL NEWS, vol. xl., p. 225), in common, I believe, with most analysts, condemns Eggertz's colorimetric method for the estimation of carbon in steel when accurate results are required, although, judging from the figures he gives, it works in his hands very fairly. M. Kern seems to favour the chromic acid method in cases where strict accuracy is essential; but I should like to ask him if he has ever tried the direct combustion of the steel in a short platinum tube in a stream of oxygen, as recommended by me in a paper read before the Chemical Society in 1870.* This method is exact and rapid, the combustion of 2 grms. of steel not requiring more than forty minutes, and the quantity of ferric oxide formed is a check on the completeness of the combustion. For a steel works, where many carbon estimations are required, I should imagine the method to be particularly suitable, as only the boat containing the sample need be changed, and in a series of experiments, made immediately one after the other, one weighing of the potash bulbs would suffice for each estimation after the first.

The chromic acid method as modified by Elliott, involving filtration through asbestos, I found to be tedious, and liable to error.

I append some results obtained by the three processes:—

Eggertz's Method (Mean of several).

1.	2.	3.	4.	5.	6.	7.	8.
0.283	0.349	0.486	0.587	0.701	0.789	—	1.319

Chromic Acid Method.

0.349	0.421	0.457	0.670	0.724	0.900	0.943	1.248
—	0.533	0.554	—	—	0.782	0.942	—
—	—	—	—	—	0.734	—	—

Direct Combustion.

0.273	0.359	—	—	0.649	0.763	0.921	1.180
—	0.359	—	—	0.620	0.759	0.922	1.151
—	—	—	—	—	0.758	—	—

—I am, &c.,

W. DOUGLAS HERMAN.

St. Helens, Lancashire,
November 17, 1879.

BEHAVIOUR OF CHLORINE AT A HIGH TEMPERATURE.

To the Editor of the Chemical News.

SIR,—After reading the letter published in your impression of November 14 (vol. xl., p. 232) from Prof. Victor Meyer

* *Jour. Chem. Soc.* [2], viii., 375.

to Dr. Endemann, I immediately wrote to my friend, Prof. Meyer, and have just received the following reply from him, and as it is written in English, I can give it verbatim.

"Zürich, November 20, 1879.

"MY DEAR MR. WATSON SMITH,

"In reply to your kind letter of the 15th November, I beg to inform you that the following statement, made by me to Dr. Endemann and printed in the CHEMICAL NEWS of the 14th of November:—'That the communications of the English journals, which refer to the obtaining of oxygen from chlorine, were published without my knowledge and against my wish,' refers to the article 'Decomposition of Chlorine' (CHEMICAL NEWS, 15 Aug., 1879), and to the communication of Sir B. C. Brodie (*Journal of the Chemical Society*, No. 202, Sept., 1879, p. 679.) But I did not in the least have in view your article (CHEMICAL NEWS, 1st August, No. 1027, p. 49), the contents of which, as far as my researches are concerned, have met with my full approval.

"Sincerely yours,

"VICTOR MEYER."

It is possible some misunderstanding might have arisen had my position, with regard to the publication of the article under the title heading this communication, been left at all in question.—I am, &c.,

WATSON SMITH, F.C.S., F.I.C.

The Owens College, Manchester,
November 24, 1879.

Decomposition of Chlorine.—MR. FREDERICK BARKAS, of the Zurich Polytechnikum, writes as follows:—A most important chemical discovery has just been made by Herren Victor and Carl Meyer, of the Polytechnikum, Zurich. Herr Victor Meyer had been making a number of experiments to determine the vapour-densities of some organic compounds whose constituents were doubtful. Having invented a new and simple apparatus for the purpose, in order to determine its accuracy he made several experiments to test the vapour-density of the commoner elements—oxygen, &c.—at temperatures from 100° C. (boiling water) to 1567° C. (heat given by a gas furnace). At length he tried chlorine, which gas was obtained by heating pure dry bichloride of platinum, but the results were not in accordance with theory. When the gas was heated at temperatures under and up to 620° C. it gave a vapour-density 2.46, while theory gives 2.45. This was very good; but at 808° C. the density was only 2.20. At 1028° C. it gave 1.87, while at from 1242° C. to 1567° C. the density remained nearly constant at 1.64 average. From this it was to be inferred that two molecules of chlorine at temperatures above 1200° C. break up into three molecules. Next came the question, Does this arise from an alteration of the molecular constitution of chlorine, or from an actual decomposition into some new gases; in other words, is chlorine an element? Thereupon the expanded chlorine gas was slowly caused to stream into a fluid that absorbs chlorine. Potassa, iodide of potassium, and mercury were all used for the purpose, and with the same result—a gas accumulated in the measuring tube that was not chlorine, but oxygen. Chlorine was thereby proved to be not an element, but an oxide of some new element. A number of careful investigations were then made to be sure that the chlorine used was absolutely pure and dry, but with the same result. The new element, hypothetically called *Murium*, has not yet been isolated, but the learned professors are carrying out the important investigation with all diligence, so that doubtless within the course of a few weeks we shall hear more of the new element, *Murium*. That chlorine is an oxygen compound is not altogether a new idea. Sir Humphry Davy, after his celebrated discoveries of the compound nature of soda and potassa, surmised that chlorine, iodine, and bromine were likewise oxygen compounds.—*Journal of the Franklin Institute*, November.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 18, November 3, 1879.

Abnormal Spectrum of Light.—M. de Klercker.—The anomaly in the spectral position of the various luminous rays, produced when the light is dispersed by the solutions of certain colouring-matters, has for some years attracted the attention of physicists. From his experiments the author concludes that the abnormal luminous spectrum is composed of two parts entirely distinct, and due, doubtless, to the different magnitude of the retardation produced by the molecules of different kinds contained in the solution.

Stomachic and Duodenic Digestion: Action of the Pancreatin.—Th. Defresne.—Hydrochloric acid in the gastric juice is combined with an organic base, which moderates its action and changes its properties. Under this influence pepsic digestion is comparable to that which takes place in the stomach. The acidity of the mixed gastric juice, after half an hour of ingestion, is not due to leucin hydrochlorate, but to the lactic, sarcolactic, tartaric, and malic acids.

Ultra-violet Absorption Spectra of the Nitric and Nitrous Ethers.—J. L. Soret and A. A. Rilliet.—Ethyl, butyl, and amyl nitrates absorb energetically the ultra-violet rays. The action of amyl and ethyl nitrites is very similar. The spectrum of these ethers in an alcoholic solution is particularly interesting in the solar light.

A New Stellar Spectroscope.—L. Thollon.—Not susceptible of useful abstraction.

Vapour-tension of Saline Solutions.—F. Pauchon. The value of α (one of the coefficients drawn from the author's experiments) in case of potassium and sodium chlorides and sodium nitrate and sulphate increase with the quantity of salt dissolved, whilst in case of sodium nitrate and sulphate it increases.

An Electro-capillary Thermometer.—E. Debrun.—The principle of the apparatus is as follows:—In a Lippmann's electrometer any mechanical action which modifies the form of the mercurial meniscus determines an electric action capable of giving rise to a current whose force is proportional to the mechanical action exerted.

Animal Cellulose or Tunicin.—M. Franchimont.—The difference between animal cellulose and that of plants, if it exists, cannot be attributed to a difference of the groups $C_6H_{10}O_5$. It depends upon a different degree of polymerisation, or a more intimate isomerism.

Researches on the Different Modes of Combination of Phosphoric Acid in the Nervous Substance.—L. Jolly.—Phosphoric acid exists in a phospho-conjugated state, the acid becoming free after ignition and in combination with potassium, calcium, magnesium, and iron.

No. 19, November 10, 1879.

The Thermic Absorbent and Emissive Powers of Flames and the Temperature of the Voltaic Arc.—F. Rosetti.—The transference of flames is very great, and they exert a very feeble absorption upon the thermic radiation by which they are traversed. The transference diminishes and the absorption increases proportionally if the thickness of the flame increases. The absolute thermic emissive power of white flames produced by coal-gas is equal to unity, whilst the absolute thermic emissive power of the pale blue flames of a Bunsen burner may be expressed by 0.3219. The voltaic arc has a much lower absorptive power. The maximum intensity for the positive extremity of the carbon is about 3900°, that of

the negative extremity being only 3150°. The temperature of the arc between these two extremities is about 4800°, whatever may be the thickness of the arc and the intensity of the current.

Researches on the Passivity of Iron.—L. Varenne.—The author has observed that iron, rendered passive, if subsequently plunged into ordinary nitric acid and struck against the sides of the containing vessel, is dissolved at once, the weaker the acid the smaller is the force of the blow required. Passive iron is completely immersed in dilute acid, and then removed with caution so that it is completely covered with the acid and suspended in the air. After a few moments, almost immediately, in a brisk current of air the attack begins and is continued with energy. If passive iron is plunged into dilute nitric acid, and some bubbles of air are allowed to pass near the metal, the reaction begins. The iron during its passive state is coated with a gaseous covering, which appears to consist of nitrogen binoxide. The author purposes the further investigation of these phenomena.

Alcoholic Fermentation.—M. Cochin.—The author concludes, in opposition to M. Berthelot, that yeast does not give rise to a soluble alcoholic ferment.

Additional Note on the Calcination of the Dregs of Beet-root.—C. Vincent.—The author contravenes the charges urged against his process by MM. Du villier and Buisine (*Comptes Rendus*, lxxxix., p. 48, 1879).

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 12, 1879.

Synthesis of Phenyl-naphthalin.—Watson Smith.—A preliminary communication. The author passed a mixture of mono-brom-naphthalin and benzol both through an empty ignited tube and through one filled with soda-lime. In the second case there were formed some diphenyl and a considerable quantity of naphthalin. In the first case there appeared a little resinous matter, and an escape of hydrobromic acid gas.

Action of Oxalic Acid upon Carbazol.—W. Saida.—The product is a blue substance, soluble in cold concentrated sulphuric acid with a pure blue colour. The original compound is not separated out on the addition of water. The authors have applied to their product the name ortho-amido-phenyl-benzoic acid.

Hydratation of Terpen.—F. Flawitzky.—A preliminary communication.

Salt of Roussin.—O. Pawel.—The author considers that for this salt four rational formulæ are possible. The different statements concerning its composition are due to the fact that Roussin, Rosenberg, and Demel examined an impure decomposed mixture of potassium and ammonium salt.

Six Lecture Experiments.—C. von Thann.—These experiments cannot be intelligibly described without the accompanying engravings.

Constitution of Alizarin Blue.—C. Graebe.—The author considers that the formation of alizarin blue depends on the production of quinolin.

Action of Ammonia upon Sulpho-anthraquinonic Acids.—R. Bourcart.—The author obtained a compound corresponding in composition to that of an amido-oxo-anthraquinone. Bisulpho-anthraquinonate of soda, resembling iso-purpurin, if heated with ammonia, yields a nitrogenous acid.

Action of Antimony Trichloride and Bismuth Trichloride upon Aromatic Hydrocarbons: Characteristic Colour Reactions.—Watson Smith.—With antimony trichloride chemically pure naphthalin gives no colouration; if traces of impurities are present a crimson colouration is produced. Anthracen added in minute traces to the melted trichloride, gives a greenish yellow colour. On cooling there are formed colourless needles of

an addition product of anthracen and antimony trichloride, which appears to be characteristic of anthracen. Phenanthren dissolves in the chloride less readily than anthracen, and gives a faint greenish colour. Diphenyl and the three isomeric dinaphthyls give no colour. Stilben if mixed with the chloride at a very gentle heat gives an orange colour, which disappears if more strongly heated. The author has examined the reactions of many other bases, natural and artificial, both with antimony and bismuth trichloride.

Congeeing Point of Bromine.—J. Philipp.—Pure bromine solidifies between -7.2° and -7.3° .

New Method of Preparing Sulphodilactic Acid.—C. Böttinger.—The author obtains this acid by the action of hydrogen sulphide upon potassium pyruvate.

Behaviour of Chlorine at High Temperatures.—V. and C. Meyer.—The molecules of oxygen, nitrogen, and sulphur at 1567° still retain the molecules commonly ascribed to them, O_2 , N_2 , S_2 . The dissociation of chlorine begins somewhat above 620° ; between 800° and 1000° intermediate numbers are obtained, but from 1200° upwards the density becomes constant again, amounting to exactly two-thirds of the value calculated for Cl_2 . It is therefore proved that the molecular weight of chlorine, which is 71 up to 600° , above 1200° is reduced to 47.3 . Iodine shows similar phenomena.

On a Base Discovered in Fusel Oil.—H. Schrötter.—The body in question is a liquid which distils over between 180° and 230° , but appears to be a mixture of at least two compounds.

Oxidation Products from Cymol-sulphide.—L. B. Hall and Ira Remsen.—An examination of the derivatives of sulph-amin-para-toluylic acid.

Anhydro-sulph-amin-isophthalic Acid.—Ira Remsen and R. D. Coale.—The authors conclude that the sulph-amin group and the carboxyl group cannot co-exist in the same ortho position.

Para-oxyphenyl-acetic Acid.—H. Salkowski.—This acid, $C_8H_8O_2$, is produced during the putrefaction of horny matter. The author describes its reactions and some of its salts.

Abietic Acid.—O. Emmerling.—After a historical introduction the author describes the behaviour of this acid with bromine, zinc chloride, hydriodic acid, and with oxidising agents.

Remarks on M. Demel's Memoir on the Arseniates of Zinc and Cadmium.—H. Salkowski.—The author considers that M. Demel is acquainted with an earlier memoir of his on the same subject.

Carbonic Acid Derivatives of Isomeric Toluydins.—J. Cesack.—A brief notice of para-tolyl, meta-tolyl, and meta-ditolyl urea, and of ortho-tolyl-urethan.

Formation of Hydro-para-cumaric Acid from Tyrosin.—E. Baumann.—An account of the decomposition-products of tyrosin, and of the properties of hydro-para-cumaric acid.

Composition of "Weldon Mud" and Similar Compounds.—J. Post.—The proportion of manganese dioxide to calcium, potassium, &c. ("basis," as Weldon terms it), is much less favourable than would appear from Weldon's and Gorgeu's statements.

Influence of the Nitro and Amido Groups upon a Sulpho Group Newly Introduced into the Benzol Molecule.—J. Post.—Not susceptible of useful abstraction.

Compounds of Benzo-trichloride with Phenols and Tertiary Aromatic Bases.—O. Dœbner.—The author describes benzaurin, a colouring-matter prepared from benzo-trichloride and phenol; its reduction to di-oxy-triphenyl-methan, its combination with acetic anhydride, and its decomposition by melting potassa.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 9, October 30, 1879.

M. le Comte du Moncel, referring to the announcement of M. Peltat's researches on the action of sunlight upon galvanic batteries, points out that similar results were obtained by himself (*Comptes Rendus*, Nov. 4, 1872).

A certain M. X., suffering from asthma, is said to have used carbonic oxide with good effect to alleviate the paroxysms.

Theory of Sleep.—E. Fournié.

Practical Means of Ascertaining and Preserving the Good Qualities of Meat.—The evils resulting from an imperfect extraction of blood from the carcasses of slaughtered animals is pointed out.

Chemical Researches on Ligneous Papilionaceæ.—P. Fliche and L. Grandeau.—Already noticed.

Telephone of M. Gower.—From an English source.

Application of Electricity in Subduing Vicious Horses.—M. Sidot.

Treatment of Sewage.—M. Coquerel.—The sewage is mixed with a reagent—not specified,—the solids are separated in the form of cakes by means of a filter-press, and the press-liquor is treated with lime in order to obtain ammonia.

Quantitative Separation of Manganese from Iron.—MM. Beilstein and Jawein.—From the *Berichte der Deutsch. Chem. Gesells.*

New Method of Producing Hyponitrous Acid and Hydroxylamin.—W. Zorn.—From the same source.

New Organic Acid Contained in Agaricus Integer.—W. Thörner.—From the same source.

Purification of Water, holding in Suspension Organic Matter.—K. and Th. Müller.—This invention, secured by a German patent, consists in the use of milk of lime, followed up by carbonic acid.

Surgical Use of Compressed Air.—An anæsthetic process depending on the use of nitrous oxide mixed with oxygen and applied under pressure.

No. 10, November 6, 1879.

Agricultural operations have been carried on in the park of Noisiel by electric power.

The True Theory of the Phenomena of Interference, of Fresnel.—H. F. Weber.

Measurement of the Focal Distances of Strongly Convex Lenses.—J. Oudemans.—Mathematical papers not susceptible of useful abstraction.

Nickel Baths for Galvano-plastic.—E. Weston.—The addition of boracic acid, free or combined, prevents the formation of a subsalt of nickel at the cathode.

Oxyhydric Lamp of M. Dubosc.—The oxygen is allowed to escape from three apertures instead of one only. The apparatus is adapted for lecture rooms, theatres, &c.

Division of the Electric Light.—M. Lontin.—It seems to be believed that the electric light loses much by being divided. The author gives a case where the totalities of light obtained from 3 burners = 261 carrels; from 4 = 316 ; from 5, 305 ; from 6, 414 ; after which a regular decrease was observed.

Opposition between the Scholastic System and Modern Chemistry as regards the Chemical Constitution of Bodies.—The first portion of a lengthy memoir, in which the author seems to aim at the rehabilitation of the scholastic philosophy.

Chemical Society Research Fund.—Dr. Warren De la Rue, E.R.S., has just sent to the above fund a third donation of £100, the whole amount to be devoted to a single research.

NOTES AND QUERIES.

Action of Liquor Ammonia on Brass.—In the *CHEMICAL NEWS*, vol. xl., p. 236, I see a letter from Mr. John Y. McLellan, in which he mentions some observations he has noticed as to the action of liquor ammonia on brass, and which he implies are not as yet on record. But on referring to Vol. v. of "Gmelin's Chemistry," he will find that "Cupreous Oxide with Ammonia" is described as a colourless liquid which gradually turns blue on exposure to the air. This absorption of oxygen, by the above-mentioned, has, as is well known, been employed for the rapid estimation of oxygen by the inventor of that very convenient apparatus sold by Messrs. Mawson and Swan, of Newcastle-on-Tyne.—ARTHUR HABERSHON.

MEETINGS FOR THE WEEK.

MONDAY, Dec. 1st.—Medical, 8.30.
— Royal, 4. (Anniversary.)
— Royal Institution, 5. General Monthly Meeting.
— London Institution, 5.
— Society of Arts, 8. (Cantor Lecture). "Chemistry of Bread and Bread Making," Dr. Graham.
TUESDAY, 2nd.—Civil Engineers, 8.
— Zoological, 8.30.
WEDNESDAY, 3rd.—Society of Arts, 8.
— Meteorological, 7.
— Geological, 8.
— Pharmaceutical, 3.
THURSDAY, 4th.—Chemical, 8. Ballot for new Fellows. "On the Theory of Fractional Distillation," Part II., F. D. Brown. "On the Influence exerted upon certain Chemical Changes by Variations in the amount of Water of Dilution," M. M. P. Muir and Chas. Slater. "On α - and β -Phenanthren Carbonic Acid," Dr. Japp. "On some Derivatives of Phenyl-acetic Acid," Dr. P. Phillips Bedson.
FRIDAY, 5th.—Geologists' Association, 8.

TO INVENTORS. AND PATENTEES.

R. E. FARRANT and Co., Dextrine Manufacturers, Gorton Gum Works, Manchester, are prepared to undertake the manufacture of any Chemical or Foods Speciality, or to furnish assistance (if required) with ample and superior accommodation, with steam-power, to any inventor or patentee desiring to manufacture.

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ILLUSIONS; The Puzzled Artist, The Knight Watching his Armour, daily by Mr. J. L. King.—**THE PHYSIOSCOPE, MICROSCOPE, KALEIDOSCOPE, CHROMATOPES, &c.**—**EDISON'S LOUD SPEAKING TELEPHONE,** demonstrated by Mr. T. C. Hepworth.—The latest and most wonderful invention in DIVING, Fleuss's Apparatus, enabling the Diver to remain under water any length of time, without any assistance from or connection with the surface, demonstrated in the Large Tank by the inventor.—**THE CHEMISTRY OF COAL,** an Experimental Lecture, and FLASHING SIGNALS, by Mr. J. L. King.—**CAVES AND DEVICES,** by Mr. T. C. Hepworth.—**INSTRUMENTAL CONCERT** by the talented Mdllers, and Master Paggi.—**GRAND POPULAR VOCAL COCERT** under the direction of Mr. Steadman, every Monday at 9. Admission to the Institution 1s. Open 10 till 1, 2 till 5, and 6 till 10.

THE CHEMICAL NEWS

AND
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THE CHEMICAL NEWS.

VOL. XL. No. 1045.

THE PERIODIC LAW OF THE CHEMICAL ELEMENTS.

By D. MENDELÉEFF.

(Continued from page 256.)

I SHALL show further on, and it can be seen from the accompanying tables, that relations of this kind can be drawn up for all the elements, showing that there is an intimate dependence between their properties and their atomic weights.

We could have foreseen this by means of the atomic theory, because the atomic weight forms one of the variable magnitudes which determine the functions of atoms. A similar consideration led me to discover the above-mentioned dependence, and this is the reason why I mention it here.

From the preceding, as well as from other relations that I have succeeded in finding, it results that all the functions which show how the properties depend on the atomic weight are periodic functions. First, the properties of elements become modified as the atomic weights increase; then they repeat themselves in a new series of elements, a new period, with the same regularity as in the preceding series. The periodic law can therefore be formulated in the following manner:—*The properties of simple bodies, the constitution of their compounds, as well as the properties of these last, are periodic functions of the atomic weights of elements.*

We now pass on to the manner of finding out the function which shows us this dependence; to effect this we should begin by finding the length, or, in better words, the number of members in a period. As for expressing this function, it appears of its own accord in some cases (such as the forms of oxidation); in other cases, there are not up to the present time any means of measuring it exactly, but, nevertheless, it keeps its periodic character.

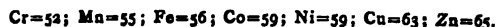
Already the above-mentioned relations have brought to light the existence and the properties of a period of seven elements, corresponding to the period Li, Be, B, C, N, O, F. Let us call it the *small period* or *small series*. If H is attributed to the first series, Li, &c., will belong to the second, Na, &c., to the third, and so on.

However, all the elements known up to the present time do not belong, as might be believed, to the little series, and, what is still more important, there exists between the corresponding members of the odd and even series (the first two excepted, as will be seen further on) a very marked difference, while the members of the odd or even series show a greater analogy between themselves. An example will prove this sufficiently.

4th series ..	K	Ca	—	Ti	V	Cr	Mn
5th series ..	Cu	Zn	—	As	Se	Br	
6th series ..	Rb	Sr	—	Zr	Nb	Mo	—
7th series ..	Ag	Cd	In	Sn	Sb	Te	I

The members of the fourth and sixth series show a greater analogy amongst themselves than they do with the members of the fifth or seventh series. Among the members of the even series there are no metalloids so marked as in the odd series; the last members of the even series resemble in many respects (as the forms of lower oxidation) the first members of the odd series. It is thus that Cr and Mn resemble Cu and Zn in their basic oxides. On the other hand, there are marked differences between the last members of the odd series (haloids), and the first members (metals of alkalis) of the even series

which follow them. But at the same time, between the last of the even series and the first of the odd are arranged in order according to their properties and atomic weights, all the elements which cannot find room in the small periods. It is in this manner that Fe, Co, and Ni in placing themselves between Cr and Mn, on the one side, and Cu and Zn, on the other, form the following series of transition:



In the same manner as Fe, Co, and Ni follow the fourth series, Ru, Rh, and Pd follow the sixth, and Os, Ir, and Pt the tenth. These two series (one even and one odd) with the intermediate series of elements, which have just been named, form a large period comprising seventeen members. As the intermediate members (such as Fe, Co, Ni) do not correspond to any of the seven groups of the small period, they form an independent group (the eighth); the members of this group—

Fe = 56;	Ni = 59;	Co = 59
Ru = 104;	Rh = 104;	Pd = 106
Os = 193?	Ir = 195?	Pt = 197

are analogous between themselves, in the same manner as the corresponding members of the even series; such as V, Nb, Ta or Cr, Mo, W, &c., This analogy is derived from the following facts:—

1st. The metals of the eighth group are all of a grey colour, and are difficultly fusible. The fusibility increases from Fe to Co and to Ni; from Ru to Rh and to Pd; from Os to Ir and to Pt.

2nd. These metals possess, even compared with the neighbouring members, very low atomic volumes; for example, the atomic volume of Cr=7.6, of Mn=7.0, of Fe=7.2, of Co=7.0, of Ni=7.0, of Cu=7.2, of Zn=9.2. The volume of Mo=11.2, whilst the volumes of Ru, Rh, and Pd are approximately 9, that of Ag is 10.3, that of Cd 13.0; the volumes of Os, Ir, and Pt are about 9.5, and of W 10.1; that of Au is 10; and, lastly, that of Hg is 15. The smallness of volumes, or of the distances between the atomic centres, render the metals of the eighth group difficultly fusible, leaves them only a small amount of chemical energy, and also determines other properties.

3rd. These metals possess in the highest degree the power of condensing and abandoning oxygen, as has been shown for Ni, Pd, Fe, and Pt by Graham and Raoult.

4th. Their highest forms of oxidation are either bases or acids of such a feeble character that they are easily transformed into lower oxides with a more marked basic character.

5th. In this group we only meet with metals whose form of combination is RO_4 or R_2O_8 , such as OsO_4 and RuO_4 ;* (it is for this reason that these metals have been collected together in a special group.) We notice that in each series the highest oxides capable of being formed by the metals from Fe to Cu, from Ru to Ag, from Os to Au, contain less and less oxygen. The highest oxide of iron is FeO_3 ; of cobalt, CoO_2 ; of nickel, Ni_2O_3 . In the same manner, osmium gives us OsO_4 ; iridium with difficulty forms IrO_3 ; platinum will only give PtO_2 ; and gold, Au_2O_3 .

6th. They give stable alkaline double cyanides. Fe, Ru, and Os give combinations analogous to K_2RCy_6 ; Co, Rh, and Ir form salts whose composition is according to the formula K_3RCy_6 ; Ni, Pd, and Pt give salts according to the general formula K_2RCy_4 .

7th. They form stable ammoniacal compounds, and resemble one another in many respects. For example, Claus has prepared salts of rhodium and of iridium, which correspond to the roseo-salts of cobalt, $\text{RX}_3 \cdot 5\text{NH}_3$, such as $\text{RhCl}_3 \cdot 5\text{NH}_3$.

8th. Some forms of combination of these metals, particularly the higher forms, are distinguished by their characteristic colours, &c.

* Ferric acid would apparently be FeO_4 ?

TABLE I.

TYPICAL ELEMENTS.			LARGE PERIODS.				
H = 1	Li = 7	Na = 23	K = 39	Rb = 85	Cs = 133	"	"
B = 11	Be = 9.4	Mg = 24	Ca = 40	Sr = 87	Ba = 137	"	"
C = 12	B = 11	Al = 27.3	"	? Yt = 88?	? Di = 138?	Er = 178?	"
N = 14	C = 12	Si = 28	Ti = 48?	Zr = 90	Ce = 140?	La = 180?	Th = 231
O = 16	N = 14	P = 31	V = 51	Nb = 94	"	Fa = 182	"
F = 19	O = 16	S = 32	Cr = 52	Mo = 96	"	W = 184	Ur = 240
	F = 19	Cl = 35.5	Mn = 55	"	"	"	"
			Fe = 56	Ru = 104	"	Os = 195?	"
			Co = 59	Rh = 104	"	Ir = 197	"
			Ni = 59	Pd = 106	"	Pt = 198?	"
			Cu = 63	Ag = 108	"	Au = 199?	"
			Zn = 65	Cd = 112	"	Hg = 200	"
			"	In = 113	"	Tl = 204	"
			"	Sn = 118	"	Pb = 207	"
			As = 75	Sb = 122	"	Bi = 208	"
			Se = 78	Te = 125?	"	"	"
			Br = 80	I = 127	"	"	"

TABLE II.

SERIES.	GROUP I.	GROUP II.	GROUP III.	GROUP IV.	GROUP V.	GROUP VI.	GROUP VII.	GROUP VIII.
	R ₂ O	RO	R ₂ O ₃	RH ₄ RO ₂	RH ₃ R ₂ O ₅	RH ₂ RO ₃	RH R ₂ O ₇	RO ₄
1	H=1	"	"	"	"	"	"	"
2	Li=7	Be=9.4	B=11	C=12	N=14	O=16	F=19	"
3	Na=23	Mg=24	Al=27.3	Si=28	P=31	S=32	Cl=35.5	"
4	K=39	Ca=40	--=44	Ti=48	V=51	Cr=52	Mn=55	Fe=56; Co=59 Ni=59; Cu=63.
5	(Cu=63)	Zn=65	--=68	--=72	As=75	Se=78	Br=80	"
6	Rb=85	Sr=87	?Yt=88	Zr=90	Nb=94	Mo=96	--=100	Ru=104; Rh=104 Pd=106; Ag=108.
	(Ag=108)	Cd=112	In=113	Sn=118	Sb=122	Te=125	I=127	"
	Cs=133	Ba=137	?Di=138	?Ce=140	"	"	"	"
	"	"	"	"	"	"	"	"
	"	"	?Er=178	?La=180	Ta=182	W=184	"	Os=195; Ir=197; Pt=198; Au=199.
	(Au=199)	Hg=200	Tl=204	Pb=207	Bi=208	"	"	"
12	"	"	"	Th=231	"	Ur=240	"	"

Above are two tables in which will be found supplementary particulars about what has been already said. In the first the elements are placed in *large periods*, with their atomic weights. In the second they are arranged in groups and series, that is to say, in *small periods*, in such a manner that the differences between the odd and even series become very apparent.

Observations on Table I.—For the sake of brevity the atomic weights have been given in these tables in round numbers, because in the greater number of cases we cannot be sure of the exactitude of the tenths nor of the units. A point of interrogation (?) *before* the symbol of an element means that the incomplete state of the researches on that element do not allow us yet to give it a determinate position in the system. A point of interrogation *after* the atomic weight indicates that in respect to the atomic weight of that element we are still in doubt; or, in other words, the equivalent of the element does not appear to have been fixed exactly up to the present day. Some atomic weights have been modified in the table according to the periodic law (see chapter 5). Thus the atomic weight of tellurium is put at 125? which is in accordance with the periodic law, and not 128, as found by Berzelius and others.

Observations on Table II.—In this table the groups are indicated by Roman figures. The seven first groups correspond to the seven members of each series; the eighth group has been already characterised (see above). Cu, Ag, and Au have been placed in the eighth group, because of their analogous properties; they could also have been put in the first group, because of their forms of power oxidation. The first two series have been separated

from the others, for reasons which will be explained later on. They are *typical series*.

(To be continued.)

ON A NEW METHOD FOR THE SEPARATION OF NICKEL AND COBALT.

By M. PH. DIRVELL.

THIS method is founded on the following facts:—

1. If there be added to the aqueous solution of cobalt nitrate or sulphate, an excess of a cold saturated solution of phosphorus salt, mixed with a solution of ammonium bicarbonate from which no ammoniacal odour is emitted, a bluish precipitate is formed in the liquid. When the mixture is slowly warmed the equivalent of carbonic acid in excess at first escapes; then, by boiling for a few seconds, a very distinct ammoniacal odour is perceptible. At this moment the boiling is discontinued, and from 2 to 3 c.c. of ammonia added to the liquid. The precipitate is, for the most part, dissolved, and it is only necessary to warm gently to 100° to obtain a precipitate of a beautiful purple inclining to violet, which is rapidly deposited. Analysis assigns to this precipitate the formula $\text{NH}_4\text{O}, 2\text{CoO}, \text{PO}_5 + 2\text{H}_2\text{O}$. It loses no ammonia at 110°, and is transformed at a red heat into pyrophosphate, $2\text{CoO}, \text{PO}_5$.

2. A solution of the corresponding salts of nickel, treated in the same manner, gives only a pure blue liquid, which is not rendered turbid by heat.

3. By mixing the two above-mentioned reagents in excess with a solution containing cobalt and nickel, and treating in the same manner, the red precipitate of ammonio-cobaltic phosphate is again obtained, whilst the remaining blue liquor contains the whole of the nickel. By this means the cobalt in the nickel sulphate of commerce can be detected.

I find it better to make this separation for the qualitative research of the two metals in assay flasks, where evaporation is slow. For the quantitative separation I prepare my reagents in the following manner:—1.30 grms. of phosphorus salt are digested in the cold in 250 grms. of water; 2.30 grms. of effloresced carbonate of ammonia are dissolved in the same quantity of water, and the solution saturated with carbonic acid till all ammoniacal smell is gone.

After having separated the two oxides in the customary way, and having reduced them by hydrogen, they are weighed, dissolved in nitric acid, and the acid solution evaporated to dryness, in a water-bath. The residue is re-dissolved in about 50 c.c. of water, to which is added a quantity of phosphorus salt equal to thirty times the weight of the two metals, and to which a volume of ammonium bicarbonate equal to that occupied by the phosphate has previously been added. We then operate as has already been indicated, care being taken to frequently shake the flask containing the liquid, especially after the addition of the ammonia.

If the boiling has inadvertently been too prolonged, causing the evaporation of the blue liquid containing the nickel on the sides of the vessel, and the consequent precipitation of a little nickel, it is easily ascertained by the colour of the cobalt precipitate, which will, in this case, be paler. It can also be compared with the moist ammonio-cobaltic phosphate, which is kept in a bottle to act as a test. In this case the clear blue liquor is drawn off, the red precipitate dissolved in only as much dilute phosphoric acid as is absolutely necessary (it is even better to leave a little precipitate undissolved), then the operation is continued with ammonium bicarbonate and ammonia.

In all cases the precipitate is washed in cold water, weighed on a tared filter at 100°, or calcined; 100 parts of the calcined precipitate contain 40.4 of cobalt. With regard to the blue liquid separated by filtration, sulphuretted hydrogen completely precipitates the nickel. The precipitate is calcined in a crucible with some sulphur and weighed in the state of sulphide.

This method is an exceedingly rapid one, for the separation requires only one or two hours at the most.*

In conclusion I would say that nickel sulphate now prepared with garnierite is found to contain magnesia, which affects the cobalt precipitate if the separation from that salt is attempted.—*Comptes Rendus*, November 24, 1879.

Production of Nitrous Acid on the Neutralisation of Opposite Electricities.—Prof. Böttger.—This phenomenon ensues both in dry and in moist air. It was formerly supposed that in the latter nitric acid alone was formed.—*Polyt. Notizblatt*, xxxiv., 334.

* M. Pisani, in whose laboratory this work was done, was kind enough to submit it to some analytical experiments, and he discovered that ammonium acetate could be substituted for the bicarbonate. The acetate is very simply prepared by saturating acetic acid at 8° with ammonia. Only 2 c.c. of this acetate is added, 0.050 grms. of cobalt, and 5 c.c. of the solution of microcosmic salt, prepared in the manner already indicated. In order to estimate the various quantities of cobalt in the solutions containing the two metals we rely on the following:—A rose solution contains an excess of cobalt as compared with nickel; a brown solution, half of cobalt and half of nickel; a dirty green liquid, 1 of cobalt to 3 of nickel; and a green solution, 1 of cobalt to 4 of nickel. In the first case, the acetate and phosphate are added as if all was nickel; in the last as if the liquid contained only a fourth part of nickel. After having heated the mixture of metals, phosphate and acetate, for a few moments in the water-bath, it is mixed with a little ammonia, and again heated in the water-bath. At the end of a quarter of an hour the cobalt is precipitated. In cases where the precipitate has not the desired tint it is again treated as indicated above.

RESEARCHES ON THE ACTION OF ORGANIC SUBSTANCES ON THE ULTRA-VIOLET RAYS OF THE SPECTRUM.

PART III. ON EXAMINATION OF ESSENTIAL OILS.*

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Associate of the Royal School of Mines.

MUCH chemical and physical research by various investigators has been devoted to the class of bodies known as Essential Oils; as, for instance, the work of Dr. J. H. Gladstone (*Journal of the Chemical Society*, vol. xviii., p. 1; vol. xxiii., p. 147; vol. xxv., p. 1); of Dr. C. R. A. Wright (*Journ. Chem. Soc.*, vol. xxvi., pp. 549 and 686; vol. xxvii., pp. 1, 317, and 619, "Isomeric Terpenes and their Derivatives"); and of Dr. W. A. Tilden (*loc. cit.*, vol. xxviii., pp. 514 and 1258); as well as of many others.

The new method of research employed by us, and described in a Paper about to be published in the *Philosophical Transactions*, has been applied to the examination of these substances. We have to acknowledge the kindness with which several gentlemen have supplied us with samples of essential oils, namely, Dr. Gladstone, Mr. Farries (of the firm of Burgoyne, Burbidges, Cyriax, and Farries), Dr. Septimus Piesse, and Dr. W. A. Tilden.

As in our previous experiments (Abstracts of Parts I. and II. *Proc. Roy. Soc.*, No. 192, 1879), photographs were taken of the spectrum transmitted by the undiluted liquid, and then of that transmitted by the liquid in various states of dilution, the dilutions ranging in some cases from 1 in 50 to 1 in 500,000 volumes of alcohol.

The following is a list of substances examined, classified according to the optical properties they were found to possess:—

Oils and Hydrocarbons transmitting Continuous Spectra

Australene, from oil of tur-	Juniper.
pentine.	Lavender.
Birch bark.	Lign-Aloe.
Cajputene dihydrate.	Melaleuca Ericifolia.
Caraway hydrocarbon,	Menthol, from oil of mint.
(No. 2).	Nutmeg hydrocarbon.
Calamus.	Patchouli, oil of (Nos. 1
Citron.	and 2).
Citronella.	Rose, otto of.
Cedar wood.	Rosewood.
Cedrat hydrocarbon.	Rosemary.
Cubeba.	Santal wood.
Elder.	Terebene.
Hesperidene, from oil of	Terebenthene.
orange-peel.	Vitiver.
Indian Geranium.	

Hydrocarbons showing the Absorption-Bands of Cymene.

Thyme.	Nutmeg.
Lemon.	Caraway (No. 1).

Substances showing Strong Bands of Absorption in the Spectrum transmitted by Dilute Solutions.

Oils of Aniseed.	Oils of Pimento.
" Bay.	" Thyme.
" Bergamot.	Carvol, oxidised derivative
" Bitter Almonds.	of caraway oil.
" Cassia.	Myristic, the same from
" Cloves.	nutmeg oil.
" Peppermint.	Patchouli, blue oil of.

For the most part these latter substances are known to contain the aromatic nucleus as an essential part of their

* Abstract of Paper read before the Royal Society, November 20 1879.

constitution. Thus the oils of bay, pimento, and cloves contain the substance eugenol, or—



oil of cassia consists of cinnamic aldehyd, —



oil of aniseed of anethol, $\text{C}_6\text{H}_4\cdot\text{OCH}_3\cdot\text{C}_3\text{H}_5$; and oil of thyme contains thymol, $\text{C}_6\text{H}_3\cdot\text{OH}\cdot\text{CH}_3\cdot\text{C}_3\text{H}_7$, as well as much cymene, $\text{C}_6\text{H}_4\cdot\text{CH}_3\cdot\text{C}_3\text{H}_7$.

Some other oils, such as bergamot and oil of peppermint, as likewise the bodies menthol, carvol, and myristicol, have an unknown constitution. The three latter substances are known to be isomeric (*Journ. Chem. Soc.*, vol. xxv., p. 1).

Great interest is attached to our examination of these bodies, since we consider it to be proved from the character of the spectra they transmit that the nucleus of menthol is a terpene, while the benzene ring is the inner basis of carvol and myristicol. Bergamot appears to be a terpene mixed with some derivative of the aromatic series; but the oil of peppermint, on the other hand, is essentially a substance belonging to this latter class. The refraction equivalents of carvol and myristicol are abnormal, like those of benzene derivatives, a fact which confirms our conclusions regarding the constitution of those substances.

The following is a summary of our observations with regard to the terpenes:—

1. The terpenes with the composition $\text{C}_{10}\text{H}_{16}$ possess, in a high degree, the power of absorbing the ultra-violet rays of the spectrum, though they are inferior in this respect to benzene and its derivatives, to which class of bodies they are so closely allied.
2. Terpenes with the composition $\text{C}_{15}\text{H}_{24}$ have a greatly increased absorptive power for the more refrangible rays,—that is to say, they withstand dilution to a greater extent the greater the number of carbon atoms in the molecule.
3. Neither the terpenes themselves nor the oxidised or hydrated derivatives occasion absorption-bands under any circumstances when pure, but always transmit continuous spectra.
4. Isomeric terpenes transmit spectra which generally differ from one another in length, or show variations on dilution.
5. The process of diluting with alcohol enables the presence of bodies of the aromatic series to be detected in essential oils, and even in some cases the amount of these substances present may be estimated.

Several diagrams in illustration of the kind of absorption excited by the different substances are presented with the complete Paper.

ON THE USE OF THE POLARISCOPE IN DETERMINING THE PROGRESS AND COMPLETION OF OXIDATION IN CONVERTING ANTHRACENE, CRUDE OR OTHERWISE, INTO ANTHRAQUINONE.

By BENJ. NICKELS, F.C.S., F.I.C.

It is well known that the crystalline form assumed by anthracene and its oxidised product anthraquinone differ widely, the former as deposited from a tolerably strong benzol solution, in sharp tabular overlaying plates; anthraquinone, on the other hand, in distinct needle and stellated groups.

Examined polariscopically, both compounds present very beautiful objects, the former as crystallised in the

tabular form exhibiting a superb play of colours; while from a weaker solution, small, but on characteristic leaflets of an intense cobalt blue tint, anthraquinone similarly viewed presenting coloured bands only, crossing the needles individually or grouped.

Other substances accompanying crude anthracene, such as carbazol, acridine, phenanthrene, pyrene, chrysene, also exhibit to an extent distinctive and characteristic forms, but as compared with anthracene, whether as hydrocarbons or oxidised products, so entirely opposite that limited observation readily distinguishes them. Naphthalene is the only body in any way resembling anthracene, but here again with equally characteristic difference, and to a careful observer there need be no error in judgment.

As will readily be seen, mixtures of these substances should be distinguishable with the same facility as the starches, and to an extent this assertion is borne out, but not without some care and experience in the preparation of the sample to be examined. It is, however, in admixture with oxidised, or partially oxidised anthracene, that the mode of observation indicated becomes of value, and a very ready method of following the process of oxidation as it proceeds. This operation, thanks to the researches and elaborate publications of Perkin, Caro, Graebe, Liebermann, and others, is now so well known that little need be said concerning it. Briefly, however, partially purified anthracene is subjected to the action of boiling potassium bichromate and diluted sulphuric acid. When employed in carefully adjusted quantity the anthracene only is attacked, and the associated or unoxidised bodies afterwards removed by further processes also equally well understood.

The operation is one of some nicety, inasmuch as too little "bichrome" involves actual loss of anthracene, and too much, complicating after process in purifying the resulting anthraquinone.

In view of the necessity of this close adjustment in the use of oxidising agents, it has been suggested to make laboratory trials on each and every sample of crude material before passing it into the works. Without in any way denying or calling in question the value of this mode of examination, I would venture to think the operation may be more easily controlled during the factory process of oxidising, and by withdrawing from time to time, as the operation proceeds, about one gramme of the partially converted anthracene from the oxidiser, washing, drying, dissolving in boiling benzol, and depositing one drop of the filtered solution on a glass slide for microscopic and polariscopic examination.

A detailed description of the varying appearance, as the operation proceeds, would involve much writing, but sufficient to say it is so well marked and defined that any observer accustomed to microscopy will readily discern features not to be disregarded in proportion as the oxidation proceeds; the needles of anthraquinone become more and more defined, increasing in quantity until finally, in a well-conducted operation, they occupy the entire field of view, traces of anthracene crystallise out quite independently, and creeping towards the outer edge of the circle are at once recognised by an experienced observer. Larger proportions stud the field in the characteristic and easily traced blue leaflets. 5 per cent of anthracene left unoxidised in the finished product for "crude," and as indicated by appropriate and well-known tests is sharply and unmistakeably discriminated as anthracene, even in the presence of associated bodies, and in the course of a few minutes only, thereby enabling the manufacturer to add further quantities of oxidising material up to the exact point of final conversion of available anthracene.

I need scarcely say that some experience is necessary with this test, simple as it may appear. I shall have much pleasure in exhibiting its details, as occasion permits, to those sufficiently interested on calling at my laboratory.

Leadenhall Street E.C. Nov 25, 1879.

NOTES OF SOME OBSERVATIONS ON
NITRIFICATION.*

By EDMUND W. DAVY, A.M., M.D.,
Professor of Forensic Medicine, Royal College of Surgeons, Ireland, &c.

A good deal of attention, on the part of chemists, has of late been given to the subject of nitrification, or the formation of nitrites and nitrates under different circumstances. This has arisen, in a great measure, from the observations of MM. Schloesing and Müntz,† which were laid before the Academy of France about two years ago. From the researches of those gentlemen, they arrived at the conclusion that nitrification was due to an organised ferment, and that it was probably the office of some of the low forms of vegetable life to produce those oxides of nitrogen under different circumstances. And the subsequent investigations of Warington, Storer, and of other chemists, would appear to go far to confirm the correctness of their theory of nitrification, at least under the conditions in which their experiments were made. Though there exists, no doubt in many cases, an intimate relation between the formation of nitrites and nitrates, and the development of certain organised germs, still as far as my observations go, I do not think that there is sufficient proof to show that their development in such instances is the cause of nitrification, and not, rather, one of the circumstances attendant on that process.

My experiments, however, were made not with a view to determine that question, but in reference to the detection of animal impurities in potable waters, and to ascertain the circumstances which were favourable or otherwise to the formation of nitrites and nitrates in waters which were so polluted, as the presence of such salts is generally regarded as indicating previous sewage contamination, and the drinking of water with such pollution is not only injurious to the health of those who thus employ it, but there exist strong grounds for the opinion which is now very generally entertained, that such water frequently becomes the means of conveying the germs of certain formidable diseases, especially those of typhoid fever and cholera, from its containing the fæcal and other emanations of individuals labouring under those maladies, and thus disease and death are often insidiously brought into many homes when such diseases are prevalent in different localities.

Besides, as the formation or production of nitrates is one of great industrial and agricultural importance, any facts which might directly or indirectly enable us to facilitate or hasten that process would be of much practical value.

As human urine and feculent matters may justly be regarded as the most offensive and dangerous ingredients of sewage in general, my experiments have been confined to those matters, and were principally made on urine, which, from its containing different nitrogenous substances, readily susceptible of decomposition, is peculiarly suited for the study of the nitrification of animal matters. By mixing this liquid with various proportions of water, and placing the mixtures under different circumstances, I have endeavoured to ascertain those that were favourable or otherwise to their nitrification; and to determine some points connected with that process which required further investigation. I should here observe that in detecting the occurrence of nitrification I have principally used the well-known test of Price for nitrites, which consists in adding to the water or mixture a thin solution of starch, containing a little iodide of potassium, and acidifying with diluted sulphuric acid, when a blue reaction from the liberated iodine will be immediately produced, should a very minute quantity of a nitrite be present. And as there is every reason to suppose that the production of

nitrites precedes that of nitrates in the nitrification of organic matters in solution, and the detection of the former is much more easily effected than the latter, at least under the conditions existing in my experiments, I was satisfied in most cases to obtain the evidence of the formation of nitrites by the employment of the test to which I have just referred.

The experiments of Warington* have led him to conclude that darkness is an essential condition to the development of those low forms of vegetable life which are supposed in many instances to give rise to nitrification.

This is a question which it is difficult to determine decisively one way or the other, owing to the impossibility of having with us continuous daylight to operate with. Still I think we may arrive at an approximative conclusion on this point, by making comparative experiments on similar mixtures, kept altogether excluded from the light, and on those exposed to its full influence, and then determining the amount of nitrification which had taken place in each, after a given time; and if darkness be so essential to that process, we should naturally expect that in the mixtures exposed to its continuous influence there would be an earlier and a greater development of nitrification, than in those which had been placed under it for about one-third or one-half the time, each day of twenty-four hours.

From the results of several comparative experiments made in this way, I have come to the conclusion that the conditions of light or darkness exercise but little influence one way or the other in this process, at least under the circumstances existing in my experiments, which consisted in placing different portions of the same mixtures in similar bottles, some of which were surrounded with black cloth or velvet to exclude light, whilst others were left uncovered, and all of them were suffered to remain open or uncorked. On examination after a few days there was but little difference as to the amount of nitrification that had taken place in each—indeed in some of my experiments it had progressed to a greater extent in the uncovered than in the covered bottles; and in all made on this subject (except those to determine this point as to the necessity or not of darkness), the mixtures were left exposed to the light, and some to the full influence of strong sunshine, yet still a considerable amount of nitrification took place in each. Besides, in nature much of the nitrates which occur in the surface soils of different localities must have been formed under the influence of more or less daylight; all of which facts, I conceive, are more or less opposed to the necessity of darkness in this process.

Another point which has not, I believe, been clearly established, at least as regards nitrification occurring in water containing organic matters, is the necessity of having a certain amount of air or free oxygen to carry on the process; this I have proved in the following very simple manner:—To water which had been kept boiling for some time to expel its contained air, I added a small quantity of freshly voided urine (the proportion employed being about one part of urine to sixteen parts of water, such a mixture having been proved to be very suitable for nitrification), and then repeated the boiling to ensure the removal, as far as possible, of any dissolved air. Several bottles which had been kept immersed in the boiling mixture were then filled completely with it, corked, and sealed with sealing-wax, to prevent the access of air. Some, however, of them containing this mixture were left open for comparison. After leaving the bottles for a day or two in the same place, I first examined the open ones for nitrites, and when the test indicated the abundant formation of those

* *Journal of the Chemical Society*, January, 1878.

† In ascertaining the amount of nitrification, the indigo process as described by Sutton in his "Volumetric Analysis" was employed which served for the determination of the nitrites and nitrates collectively; and though it may not be quite so accurate as some other methods, was sufficiently so for this purpose, as it was only the comparative amounts of nitrites and nitrates formed under the different circumstances of the experiments that I wanted to determine.

* Read before the Royal Irish Academy, May 12, 1879.
† *Comptes Rendus*, lxxiv., 301.

salts, I opened one of those sealed, when not a trace of nitrites was discoverable in its contents; the remaining sealed ones were opened at different periods subsequently, with the same results. Other comparative experiments were made, where the temperature of the mixtures was artificially kept at a heat very favourable to nitrification, but, in every instance where the access of air had been excluded, no trace of nitrites could be detected—clearly proving the necessity of more or less air or free oxygen for their formation. But the amount necessary to commence, at least, the process, is small, for I found where the mixture had not been boiled previously to the complete filling, corking, and sealing of the bottles, that the air dissolved in the liquid was sufficient to cause the production of nitrites to some extent.

The quantity of animal matter which is held in solution in the water, I find exercises a considerable influence over nitrification; for where it occurs in very large proportion, there the process either does not take place at all, or is carried on much slower than in the more dilute solutions. This I have proved by comparative experiments with water mixed with different proportions of the same sample of urine, or of solution of excrementitious matter, where I found that nitrification occurred first in the more dilute mixtures; and that where there was much organic matter present, that the nitrites which might ultimately be formed soon afterwards disappeared again by their subsequent change or decomposition, whereas those that had been produced in more dilute solutions have remained unchanged for a considerable time.

But the circumstance which I have found to exercise the greatest influence over nitrification is that of temperature; for I have observed that in cold weather it is very slow in taking place, whilst in warm it is much quicker, and that by the application of artificial heat the process can be greatly accelerated. The correctness of this observation is borne out by the well-known fact, that it is from the soils of different hot climates that we obtain our chief supply of nitrates.

As to what may be the most favourable temperature for this process, I have not yet been able to determine, owing to the difficulty, as I am circumstanced, in maintaining continuously the same degree of artificial heat; but I have found that where the mixtures were placed where they were kept at a temperature which varied from about 70° to 80° F., that there the process was carried on very quickly, and that nitrites were soon abundantly formed, whereas similar mixtures maintained at lower degrees of heat, or at the ordinary temperature, not a trace of those salts could be detected in the same time, and that their presence was not discoverable till after a much longer period.

The foregoing observations have, I conceive, some important bearings as regards the contamination of water with sewage, and the evidence of such, derivable from the occurrence in it of nitrites and nitrates. For though the presence of those salts is undoubtedly in many instances an indication of previous sewage pollution, still their absence, taken by itself, cannot be relied on as a sure indication of the freedom of the water from such contamination. For the circumstances present may have either been unfavourable to the formation of nitrites and nitrates, or have produced their subsequent rapid disappearance—thus, for instance, the lowness of the temperature of the water may have prevented their formation, or the quantity of organic matter present may have interfered with their development, or have led to their subsequent change and disappearance. Such, amongst other circumstances influencing the presence of those salts in water containing animal matters, will at once be evident; and their absence unless accompanied by other indications of purity, cannot be relied on as a proof of the freedom from such contamination.

Before I conclude, I wish to call attention to another fact, which I have noticed in connection with this subject, viz., the rapidity with which nitrites are sometimes formed

in waters contaminated with sewage impurities. This is a subject of considerable importance in an analytical point of view, as I shall endeavour briefly to explain.

It is well known by those who have analysed potable waters, that the method which chemists now principally employ to ascertain their purity or otherwise is to determine the quantity of ammonia a given amount of the water will yield on distillation, both before and after the addition of a strongly alkaline solution of permanganate of potash. The first obtained is termed the free and the second the albuminoid ammonia. The former is regarded as the representative of the nitrogenous organic matters previously existing in the water, which have undergone more or less decomposition, whilst the latter is produced by the action of the alkaline permanganate on those substances still present in the water. Consequently, the less of each that is furnished by a sample of water when so treated, the purer organically is it regarded, and the safer, other circumstances being similar, would it be for potable purposes. When lately analysing a sample of water that had been contaminated with sewage, to ascertain the amount of such pollution, which was afterwards the subject of an important legal inquiry, in my first trial I found that the water yielded a quantity of free ammonia which was equivalent to 0.970 part of a grain per gallon, but on repeating the determination a few days afterwards, it was discovered that it had fallen to 0.186 part of a grain for the same quantity of water, or to less than one-fifth of the former amount; whereas the quantity of albuminoid ammonia yielded had slightly increased. This result as to the great decrease of free ammonia, which at first rather surprised me, I ascertained was due to the formation of nitrites, which had been developed to a large extent, in so short a time, at the expense of the free ammonia. Such being the case, if the water had not been examined till the date of the second analysis, and if the nitrites had not been taken into account, this water would have been regarded as containing much less free ammonia than it did, and consequently that the previous sewage contamination was less than it really was; this point is therefore one of some analytical importance.

It is right for me to observe, in connection with this latter fact, of the decrease of free ammonia in waters by keeping, that long after I had made that observation I met with (in the *CHEMICAL NEWS*, vol. xxxv., p. 94), a letter written by Professor Pattison Muir, of Owen's College, in which he calls the attention of chemists to some observations his brother had just made in the laboratory of the University at Sydney, in which he had noticed that the amount of free and of albuminoid ammonia, as determined by Wanklyn's process, varied very considerably with the time the sample of water had been kept; but neither of those gentlemen has offered (in the letter referred to) any explanation of the fact, further than that Prof. P. Muir throws out the suggestion, in the case of the increase by keeping of the albuminoid ammonia, that possibly it might have been owing to the germs which have escaped decomposition by the permanganate undergoing a gradual decomposition in the water, and that ammonia is one of the products of this process. Be this as it may, I have satisfied myself that the loss of free ammonia is often due to the formation of nitrites or nitrates, which are very rapidly formed under different circumstances. And as regards albuminoid ammonia, the very slight increase which I observed in my experiment was, I thought, very easily accounted for by my having in the second determination carried on the process of distillation somewhat further than in the first trial, and in this way the amount might be very naturally increased.

Finally, my observations that nitrification is greatly promoted by warmth might, I conceive, admit of some practical application in the manufacture of the nitrate of potash in the artificial nitre beds, especially in those of cold countries; and I am not aware that heat has hitherto been anywhere artificially applied to hasten or promote that important manufacture.

ON THE SEPARATION OF THE HEAVY METALS OF THE AMMONIUM SULPHIDE GROUP.

By CLEMENS ZIMMERMANN.

AFTER pointing out that the usual procedures are either very tedious or questionable as to accuracy, the author gives his method for separating zinc from the other metals of the group by means of ammonium sulphocyanide. To the liquid in question, which, in addition to the zinc salt, may contain any number of the other metals of the fourth group, iron and uranium, if present, being in the state of ferric and uranic salts, he adds, if its reaction is acid, carbonate of soda till a slight turbidity appears, neutrality being the main condition for success. An excess of a solution of ammonium sulphocyanide, not too dilute, is then added, the sides of the vessel are rinsed with water at 60° to 70°, preferably by means of an Erlenmeyer flask, and a very moderate stream of sulphuretted hydrogen is introduced repeatedly, but not very long, till the odour of this gas does not disappear after the liquid has stood for some time. During the introduction of the gas the appearance of a milky-white precipitate is first perceived, and after a considerable time zinc sulphide is separated in flocks which continually become denser. The beaker is now exposed to a gentle heat till the precipitate has settled and the liquid has become clear, which may require six hours. It is then filtered, the white zinc sulphide is washed with water containing sulphuretted hydrogen and ammonium sulphocyanide, and dried. This precipitate contains all the zinc, free from the other heavy metals of the group. When dry it may be ignited in a current of hydrogen according to Rose's process (*Poggendorff*, cx., 128), or it may be dissolved in hydrochloric acid, evaporated to dryness in a weighed platinum capsule on the water-bath, mixed with an excess of elutriated mercuric oxide, pure, and free from alkali, evaporated to dryness, and ignited. Zinc oxide remains perfectly pure and without loss, and is weighed when cold.

In the filtrate from the zinc sulphide the sulphocyanides are first destroyed by means of nitric acid, which at the same time peroxidises any ferrous or uranic salts present. This operation is best effected in a roomy long-necked flask, which is heated on the water-bath, and nitric acid added by degrees in small portions until the liquid no longer becomes red, followed by decoloration. Any yellow cyanogen persulphide formed is filtered off. If the acid is added too rapidly the liquid may be projected out of the flask.

In order to separate the iron present from nickel and cobalt, the solution, which may contain ferric salts along with nickelous or cobaltous salts or both, is mixed with an excess of ammonium sulphocyanide, when the blood-red colour of iron sulphocyanide appears; a solution of "secondary" carbonate of soda is then added drop by drop, till the red colour just disappears. All the iron is thus precipitated as ferric hydroxide, none of it remaining in solution, and no cobalt or nickel being thrown down. The precipitate is allowed to settle, filtered, washed with boiling water to which a little ammonium sulphocyanide has been added, dried, ignited, and weighed. The filtrate is treated as has been directed for the filtrate from zinc sulphide. The author separates cobalt and nickel by Liebig's method with ferric oxide.

For the separation of iron and uranium the liquid, in which the metals must have been peroxidised, is brought to a boil, mixed with an excess of ammonium sulphocyanide, and aqueous carbonate of soda is added, exactly as above directed for the separation of iron from cobalt and nickel. The same process is further followed for the removal of the iron, which is found free from the slightest trace of uranic compounds.

The filtrate which contains uranic oxide in solution is first treated with nitric acid to destroy the sulphocyanogen, then neutralised with ammonia, and mixed with ammonium sulphocyanide; the precipitate of uranium oxy-sulphide

is boiled, by which it is resolved into sulphur and uranic oxide, filtered, dried, and ignited. Lastly, the uranium is either weighed as uranoso-uranic oxide, or converted into uranic oxide by very strong ignition in a current of hydrogen gas which is maintained until the product is cool.

The precipitation of uranic oxide by ammonia is greatly promoted by the addition of ammonium chloride, without which it does not take place in dilute solutions.—*Annalen der Chemie*, 199, 1.

INFLUENCE OF ACETIC ACID ON THE SEPARATION OF IRON AS A BASIC ACETATE FROM MANGANESE, ZINC, COBALT, AND NICKEL.

By JOHN JEWETT.

It has often been observed, when manganese is separated from iron by precipitating the latter as a basic ferric acetate, that some manganese is carried down with the iron precipitate. Eggertz,* calling attention to this fact, stated that this trouble could be obviated, at least to a great extent, by the presence of free acid. To this end he recommends adding to a volume of 500 c.c., after nearly neutralising with sodium carbonate, 3 c.c. of HCl (strength not given). Stöckmann communicated in *Fresenius's Zeit.* (1877, p. 172) the results of a series of experiments in separating iron from manganese in "spiegeleisen" containing about 10 per cent of manganese. He found that varying, and usually very considerable, amounts of manganese were precipitated along with the iron, in many cases equalling 10 per cent of all that was present, and concludes that it is absolutely necessary to re-dissolve and re-precipitate the iron and recover the manganese in the second filtrate. He alluded to the modification recommended by Eggertz, but did not adopt it, finding the free acid made the iron precipitate difficult to wash. C. Kræmer† was astonished at these results, and states that the only precaution necessary (besides thorough washing) is the addition of one or two drops of dilute acetic acid to the previously neutralised solution, before adding sodium acetate and boiling; he found that at most only one-tenth per cent of manganese goes down with the iron. The experience of G. Matzsurke‡ was, on the other hand, quite like that of Stöckmann. He found, contrary to Kræmer, that the addition of one or two drops of dilute acetic acid made no appreciable difference in the results.

In view of these statements concerning a much-used process, I have, with the advice of Prof. O. D. Allen, made some experiments to ascertain the influence of free acetic acid in this method of separating iron from manganese and the other metals, zinc, cobalt, and nickel. As a basis of operation, solutions of Fe_2Cl_6 containing free HCl, and of MnCl_2 , ZnCl_2 , $\text{Co}(\text{NO}_3)_2$, and $\text{Ni}(\text{NO}_3)_2$, of known strength (0.2 gr. of the metals to every 100 c.c. of solution), were made; then by uniting 100 c.c. of the iron solution with the same amount of any one of the other solutions, a mixture of the desired metals to be separated was obtained. A few preliminary trials with iron and manganese gave results in accordance with the statements of both Stöckmann and Eggertz. Presence of free acid decreased the amount of manganese in the iron precipitate, but too much prevented complete precipitation of the iron. To ascertain to what extent free acetic acid is efficient in keeping manganese—likewise zinc, nickel, and cobalt—in solution, when present in quantities not too great to prevent precipitation and washing of the iron, necessitated a series of experiments, in which the only variable factor was the acetic acid. In every case 2 grs. sodium acetate were used (an amount equal to ten times the weight of iron

* Berg- und Huettenmännische Zeitung, 1867, p. 187.

† *Fresenius's Zeitschrift*, 1877, p. 334.

‡ *Ibid.*, 1878, p. 78.

present), and the solutions were diluted to the same volume, viz., 300 c.c. before boiling to separate iron. The precipitated iron was examined after thorough washing, by re-dissolving and repeating the separation, and testing the filtrate from the iron precipitate, according to whether Mn, Zn, Co, or Ni was present, with Br, NaCO_3 , or NaOH. The results are here tabulated:—

Per cent by volume of Acetic Acid (1044 sp. gr. added).	Taken Fe=0.2 gr. Mn=0.2 " P.c. of Mn in and Filtrate.	Taken Fe=0.2 gr. Zn=0.2 " P.c. of Zn in and Filtrate.	Taken Fe=0.2 gr. Ni=0.2 " P.c. of Ni in and Filtrate.	Taken Fe=0.2 gr. Co=0.2 " P.c. of Co in and Filtrate.
0	2.788	—	—	—
1	0.697	2.849	1.888	—
2	0.585	2.046	1.770	0.590
3	0.193	—	1.652	—
4	0.154	—	1.468	—
5	trace	0	1.219	0.315
6	trace	0	—	—
7	trace	—	0.314	—

It was found that 4 per cent by volume of acetic acid, 1.044 sp. gr., has no appreciable bad effect on the separation and filtration of the iron; when 5 per cent is used its effect is sometimes apparent, but occasions no great trouble. On making some trials with over 5 per cent, iron was not always completely precipitated, and was very difficult to wash. Great care was taken in these experiments to have as little free acid as possible present besides that intentionally added. This end can best be attained by adding sodium carbonate to the cold and preferably concentrated acid solution until a slight precipitate forms, which no longer re-dissolves by shaking and allowing to stand three or four minutes, but imparts a turbidity to the deep red solution; HCl must then be added without longer delay, drop by drop, until the fluid, though still dark, becomes clear. After neutralising in this manner, the amount of acetic acid required to form the desired percentage of the final volume was added, next sodium acetate, and lastly the solution was diluted to the final volume (viz., 300 c.c.) before boiling. Long boiling of the precipitate was avoided, one or two minutes sufficing to make the basic acetate settle quickly after removing the heat. The precipitate was washed with nearly boiling water, to which 2 grs. sodium and 1 c.c. acetic acid, per litre, had been added, until AgNO_3 gave no reaction with the washings.

It may be concluded, from the results shown in the table, that by using 4 per cent by volume of acetic acid, 1.044 sp. gr., and adhering to the above precautions, a complete separation by one precipitation can be obtained of zinc, and one sufficiently accurate for most purposes of manganese; while the amount of nickel and cobalt that goes down with the iron lessens with increase of acetic acid. The following quantitative separations of iron from zinc and manganese were also made. Precipitation was effected in a volume of 300 c.c. containing 12 c.c. acetic acid, i.e., 4 per cent of this volume and 2 grs. sodium acetate.

Fe taken. Grs.	Zn taken. Grs.	Mn taken. Grs.	Fe found. Grs.	Zn found. Grs.	Mn found. Grs.
0.1966	0.1997	—	0.1995	0.1994	—
0.2000	0.2000	—	0.1998	0.1999	—
0.1996	—	0.2000	0.1983	—	0.2002
0.2000	—	0.2000	0.1993	—	0.2001

Other Variable Conditions.

A volume less in proportion to the iron present may doubtless be advantageous when large quantities of iron are to be precipitated. The writer, however, would not recommend, on account of increased difficulty of washing, less than 100 c.c. per 0.1 gr. Fe.

Experiments made by E. H. Smith in this laboratory, on the separation of iron from manganese, under exactly

* This acetic acid, by volumetric determination, was found to contain 33.16 per cent $\text{C}_2\text{H}_4\text{O}_2$.

the same conditions as described, except that double the amount of sodium acetate was used, gave essentially the same results.

He further ascertained that iron could not be precipitated in presence of a larger amount of acetic acid by increasing the amount of sodium acetate.

Since solutions containing free acid may be neutralised with ammonia or ammonium carbonate instead of sodium carbonate, when the presence of ammonium salts is not objectionable,—for example, in separating nickel from iron,—I have duplicated some of the above experiments with nickel, with no other change than the addition of ammonium chloride. The results are appended, showing the difference when NH_4Cl is present:—

Per cent by volume of Acetic Acid (1.044 sp. gr. added).	Per cent of Ni in and Filtrate.	Per cent of Ni in and Filtrate, when 5 grs. NH_4Cl were added.	Difference.
1	1.770	0.904	0.866
3	1.468	0.708	0.760
4	1.219	0.511	0.708

—American Chemical Journal.

NOTICES OF BOOKS.

Analytical Chemistry. By W. DITTMAR. Pp. 88. London and Edinburgh: W. and R. Chambers, 1879.

It constantly happens that beginners in practical qualitative analysis are disheartened by the mass, the variety, and the complexity of the directions given in the ordinary manuals. What with alternative processes, cross-references, cautions, foot notes, appendices, and the introduction of the rarer metals, the beginner is hopelessly puzzled, and abandons all attempt to learn the method of analysis in a thorough and systematic manner. For beginners and for those students who have no occasion to carry their analytical course very far, we can heartily recommend Professor Dittmar's compact, clear, and practical little book; and, as an introduction to larger and more complete works (like the "Qualitative Analysis" of Fresenius), it will prove useful, since, so far as it goes, its statements are accurate and its methods practicable. There is no attempt to shirk the real difficulties of analytical operations and explanations; only a wise judgment has been exercised in selecting what shall be given and what shall be withheld.

This book consists essentially of a series of laboratory exercises, beginning with dry-way tests, then come wet-way reactions, and then the application of the facts observed to what Prof. Dittmar calls "the ABC of metal analysis." Next we have schemes for the examination of solutions for metals and for inorganic acids, and finally some instructions as to the characteristic reactions of organic acids. An appendix, intended for medical students, relates to the alkaloids, sugars, starch, and urine.

The pages before us are not only remarkably free from errors but contain a large number of valuable hints and precautions, not usually to be found in small treatises of an elementary sort. We fail to discover any but quite insignificant corrigenda.

A Treatise on Chemistry. By Professors ROSCOE and SCHORLEMMER. Vol. II., *Metals*, Part II. London: Macmillan, 1879.

This volume fulfils and more than fulfils the promise of its two predecessors. It treats with adequate fulness and freshness of the metals of the iron, chromium, tin, antimony, and gold groups. Special attention has been paid by the authors to technical processes both in the way of verbal description and engraved illustrations. The metallurgy of iron and the manufacture of glass are justly cited

in the preface as examples of the pains that have been devoted to the due illustration of important operations, plant, and apparatus connected with chemical and metallurgical manufactures. The last part of the volume before us is occupied with three supplementary chapters of considerable scientific importance. The first of these treats of spectrum analysis, especially so far as terrestrial matter is concerned; the second discusses the natural arrangement of the elements and the periodic law of Mendeleeff, while the third is occupied with a description of the methods and apparatus by which Piçet and Cailliet liquefied the so-called permanent gases—hydrogen, oxygen, carbon monoxide, &c. We may say at once that these three supplementary chapters leave nothing to be desired, either as to completeness, accuracy, or fairness, while the illustrations which accompany them are of equal excellence with those which adorn the previous portions of the work under notice.

Nearly two dozen metallic elements with their more important compounds are described in the less than 500 pages of the present volume. But the authors have rightly assigned appropriately larger spaces to the accounts given of such important metals as iron and gold, while the rarer and obscurer metals are more briefly, yet quite sufficiently, described. Naturally, however, vanadium, on which Prof. Roscoe has so successfully laboured, comes in for a much more complete discussion than some of the other metals which must be regarded as scarce. We confess we should like to see a few of the important constants of each metal prefixed or suffixed to the descriptive accounts given, so as to offer to the reader in a compact tabular form for immediate reference and comparison the best and latest results of chemical and physical investigations. Another suggestion we venture to make has reference to the mineralogical references in the volume before us. We cannot but commend the introduction of some account of the minerals which form the raw material of all researches into metals and their compounds; but we cannot understand the peculiar spelling and nomenclature adopted for many of the species here introduced, nor can we explain the exclusion of some of the most interesting kinds: then, too, in some instances, quite inadmissible formulæ are assigned. We give a few instances of these three peculiarities, which, considering the remarkable pains bestowed upon almost all the departments of chemical knowledge covered by this treatise, are the more unaccountable.

We begin by asking why rhodochrosite should be changed to rhodocrozoite (p. 2); erythrite to erythine (p. 132); and uranocalcite to uranocalcite (p. 225)? The latter change is most unfortunate, implying, as it does, the presence of copper instead of lime in the mineral, although the formula given is that of an uranyl—calcium phosphate: by the bye, this species contains $10\text{H}_2\text{O}$, not $8\text{H}_2\text{O}$. Stiblithe for the mineral called stibiconise by Beudant, and stibiconite by Brush and Dana, is a form which does not commend itself on the score of euphony; nor is the memory of the Comte de Bournon honoured by changing bournonite into bournanite. It is a pity that a number of such small inaccuracies in names and formulæ should have crept into a work so carefully prepared as this valuable standard treatise on chemistry. We do not refer to mere misprints, like *musirum*, on p. 251, nor hæmogoblin (so curiously associated with spectra), on p. 495; such accidents will happen to the most expert and painful readers of proof.

Remarkable Behaviour of Silver Oxide.—Prof. Böttger.—If two measured parts of silver oxide, perfectly dry, are rubbed in a porcelain mortar with 1 part antimony sulphide the mixture readily takes fire. The same phenomenon occurs if silver oxide is ground along with amorphous phosphorus. If a drop of phenol is thrown upon silver oxide the latter is partially reduced, with the projection of sparks. —*Polyt. Notisblatt*, xxiv., 322.

CORRESPONDENCE.

OZONISERS.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xl., p. 246, is described, by Mr. Albert R. Leeds, various forms of ozonisers. He does not mention mine; and it is possible he may not have heard of it, notwithstanding that it was shown at a soirée of the Royal Society some thirteen or fourteen years since.

It was, as you are aware, composed of sheets of glass, coated with tin-foil, with longitudinal slips of glass, uncoated, placed along each side, thereby forming cells, and the whole placed in a wooden box coated with shellac varnish. When large quantities of ozone are required the cells can be built up to any height, so that a very large area is obtainable. The construction is exceedingly simple.

With this ozoniser I found the best result by exposing a small surface of foil—a strip about 1 inch wide—across the plates, highly charged. With a large surface of foil and the same battery power, a very small result was obtained.—I am, &c.,

EDWD. BEANES.

Moatlands, Paddock Wood, Brenchley, Kent,
November 25, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No. 20, November 17, 1879.

Temperature of the Decomposition of Vapours.—H. Sainte-Claire Deville.—The author remarks that the experiment of M. Troost (*Annales de Chimie et Physique*, Série v., Tome 12) on the hydrate of chloral and on its existence in the state of vapour, is one of those which may be called crucial, and has stung to the quick all the partisans of the atomistic chemistry. The procedures and the apparatus used by M. Wurtz in his researches on chloral hydrate are the same as those described by the author more than fifteen years ago. M. Berthelot has perceived that M. Deville has avoided the causes of error which attach to this apparatus as reproduced by M. Wurtz. A rise of temperature is an incontestable sign of the combination of two gases, but this does not hold good for vapours, especially for those which M. Wurtz has employed, those of chloral and of water at temperatures near their point of condensation.

Observations on a Note by M. D. Cochin relating to Alcoholic Fermentation.—M. Berthelot.—The author presented the views ascribed to him by M. Cochin, not as absolute opinions, but merely to explain the leading idea of the experiments of Claude Bernard. To refer a chemical metamorphosis to a vital act is not to explain it. On the contrary, all the efforts of physiological chemistry have for their object to analyse the material changes which take place in living beings, and refer them to a regular succession of chemical acts.

Observations on the Ultra-violet Limit of the Solar Spectrum at Different Altitudes.—A. Cornu.—The extension of the limit of the spectrum estimated in wave-lengths ought to be a millionth of a millimetre for every 663 metres of altitude. The rate of progression is conformable to the theoretic value deduced from the hypothesis of a homogeneous absorbent atmosphere, but on condition of choosing as numerical data those correspond-

ing to days when the atmosphere is purest. The actual extension of the spectrum in the ultra-violet direction is one unity (millionth of a millimetre) for every 900 metres of height.

Explosion of Carbonic Acid in a Coal Mine.—M. Delesse.—The explosion occurred in the coal mine of Rochelle (Gard). The following reasons are given to show that it was not occasioned by carburetted hydrogen:—The detonations were not accompanied by flame; the bodies and clothing of the victims bore no marks of burning, and powder and cartridges in the mine had not been ignited; fire-damp has never been observed in the mine, but an escape of carbonic acid has often been observed and has been combated by ventilation. The explosion would seem to have been merely the sudden liberation of an enormous volume of this gas.

Second Note on the Effects and Modus Operandi of Antiseptics: Their Action upon Pus.—MM. Gosselin and A. Bergeron.—Pus putrefies more slowly than blood; its putrefaction is retarded by incomplete occlusion. It is retarded by antiseptics, both in actual contact and at a distance. Camphoretted spirit, phenol at 1-50th, and alcohol at 86° are equally moderators of inflammation and preservatives against septicæmia.

Critical Reflections on Experiments Concerning Human Heat.—G. A. Hirn.—Not susceptible of abstraction.

Atmospheric Polarisation and the Influence which Terrestrial Magnetism may exert upon the Atmosphere.—Henri Becquerel.—The author infers from his researches the existence of a variable divergence between the plane of the sun and the plane of the polarisation of the atmosphere in any point whatever, and the manifestation of a magnetic influence of the earth upon the atmosphere.

Solar Spots and Protuberances Observed with a Spectroscope of great Dispersive Power.—L. Thollon.—This paper requires the accompanying illustrations.

On Chlorophyll.—A. Gautier.—The author has obtained chlorophyll in crystals belonging to the system of the oblique rhomboidal prism. On exposure to diffused light they turn slowly to a yellowish green, and ultimately become colourless. In its reactions and its elementary composition it shows an approximation to bilirubin. Both are soluble in ether, chloroform, petroleum, sulphide of carbon, and benzol. Both are withdrawn from the majority of these solvents by animal black. Like bilirubin, chlorophyll plays the part of a feeble acid, yielding soluble and unstable salts with the alkalis, and insoluble salts with all the other bases. It contains not a trace of iron. Its relations with bilirubin connect it necessarily with hæmatin.

Les Mondes, Revue Hebdomadaire des Sciences.

No. 10, 1879.

Unpublished Letter from C. F. Gauss to Sophia Germain.—An astronomical paper.

State of Alkaline Phosphates in Solutions.—J. van Bemmelen.

A New Salt with an Iridium Base.—K. Birnbaum.

State of Double Salts in Solutions.—P. H. B. Ingenhous.

Condensation-products of the Tertiary Aromatic Bases.—O. Fischer.—These four papers are all taken from the *Berichte der Deutschen Chemischen Gesellschaft*.

Vibratory Forms of the Bubbles of Glyceric Liquid.—C. Decharme.—Already noticed.

No. 11, November 13, 1879.

It is said that a M. Dalmas has succeeded in destroying the phylloxera by an electric process. The stems of the vines are wrapped round with thin copper wire, through

which is passed the current of a powerful battery. Both the mature insects and their eggs are completely disorganised. No details are given.

The Metric System or the Battle of the French Standards.—C. Latimer.—The author pronounces the metric system "anti-scientific and anti-human."

The Superficial Viscos of Liquids.—J. Plateau.—Not adapted for abstraction.

Reversion of Superphosphates.—M. Joulie.—Taken from the *Annales der Chemie et de Physique*.

Bone-glass.—M. Sidot.—The author proposes to make glass simply of phosphate of lime, which he considers will prove very valuable. It is not attacked by hydrofluoric acid.

No 12, November 20, 1879.

Chemical Nomenclature.—M. Dulaurier.—In the author's system every element is to be represented by a consonant, whilst the number of molecules present in any compound is to be expressed by the vowels. Carbonic acid is xeka; ammonia, zodré, &c.

The article on the opposition between modern chemistry and the scholastic system is continued, as also M. J. Plateau's paper on the superficial viscosity of liquids.

Journal de Pharmacie et de Chimie.

November, 1879.

The Ferment of Carica Papaya.—A. Wurtz and E. Bochet.—Already noticed.

Action of Hypochlorite of Lime upon the Propylic, Butylic, and Amylic Alcohols.—J. Regnault and E. Hardy.—The authors investigate what are the chlorinated compounds resulting from the treatment of each of these three alcohols in an isolated state with the hypochlorite and hydrate of lime according to the method of Soubeiran. The resistance of the alcohols to the action of chlorine is the greater according as they belong to a more advanced term of the series.

Bark of Pala Mabi.—M. Planchon.—This paper is chiefly of a botanical pharmaceutical character.

Memoir on the Tritungstates.—Jules Lefort.—The tritungstates are not very stable, a character which distinguishes them, on the one hand, from the bitungstates, and, on the other, from the quadri- or meta-tungstates. The author has discovered the neutral tungstates, bi-, tri-, and quadri-tungstates of the sesqui-oxides.

The Elimination of Bromine from Bromo-citraconic Acid, and a New Organic Acid.—E. Bourgois.—This acid differs from citraconic acid by possessing two equivalents less of hydrogen.

Oxidation of Formic and Oxalic Acids by Ammoniacal Oxide of Copper.—Paul Cazeneuve.—The product obtained is the ammoniacal carbonate of copper, similar to that obtained by Favre. The author is extending his researches to more complex acids.

Action of Digestive Ferments Employed in the Treatment of Dyspepsia.—M. Vulpian.—This paper is of a medical rather than of a chemical character.

Verhandlungen des Vereins zur Beförderung des Gewerbflusses. October, 1879.

This issue contains the new German regulations concerning the transport of explosives. The substances included are gunpowder and blasting-powder; nitroglycerin and its preparations; nitrocellulose, especially gun-cotton; explosive mixtures containing chlorates and picrates; fulminating silver and mercury and their preparations. Explosives can be sold only to persons whose good reputation is known to the dealer, or who can produce satisfactory testimonials from the local authorities.

Chemiker Zeitung.
No. 43, 1879.

The Kurtz Process for the Manufacture of Nitroglycerin.—Not suitable for abstraction.

No. 45.

H. Hugershoff, of Leipzig, has introduced crucible-tongs with porcelain blades.

No. 46.

Preparation of Litmus as an Indicator.—M. Kre'schmar.—Commercial litmus, ground very fine, is extracted with cold water till almost exhausted, and the extract is evaporated down with pure quartz-sand. During the evaporation hydrochloric acid is added till the liquid, after escape of the carbonic acid, appears strongly red. The brownish red dry powder thus obtained is ground up, and washed first with hot and then with cold water upon large smooth filters. The first washings are a dirty brown-red, and turbid. The water then passes through clear, and finally becomes a vinous red. The residue on the filter is perfectly dried on the water-bath. For use, a larger or smaller quantity is placed upon a filter, covered with hot water and a few drops of ammonia, and washed till the sand is exhausted. The filtrate is acidified with a few drops of very dilute sulphuric acid and then neutralised.

Alizarin Blue.—G. Auerbach.—The author gives analytical details in confirmation of a former paper.

Compounds obtained from Animal Tar.—Dr. H. Weidel and G. L. Ciamician.—Animal oil freed from bases contains as main products the nitriles of butyric, valerianic, capronic, capric, palmitic, and stearic acids, as well as pyrrol, homopyrrol, and dimethyl-pyrrol.

No. 47.

The Relation of the Litre to the Kilogramme.—For practical purposes the litre is defined as that space which is taken up by a kilo. of water weighed at 15° C., and at a mean barometric pressure of 760 m.m.

A New Alkalimetric Indicator.—Dr. E. Freise.—The author proposes an extract made by digesting 50 parts of the best rasped logwood with 1000 parts of distilled water at 40° for a day, and filtering rapidly.

C. H. Wolff's Colorimeter.—The principle of this instrument is the production of an increase or decrease in the depth of the stratum of liquid under examination until it appears equal in intensity to a standard.

Chemical Nomenclature.—G. Auerbach.—The author complains that one and the same name is often applied to two or more distinct chemical compounds. He asks also by what right is the name cœrulein given to a dye which dissolves in alkalies with a green colour, and which dyes mordanted tissues a beautiful green.

New Tanning Materials.—Dr. A. Schottky.—The author calls attention to the bark of the red and the white mangrove, and especially to that of the mahogany tree, which contains from 28 to 32 per cent of tannin.

La Correspondance Scientifique.
October 14, 1879.

On Telegraphic Lightning Conductors.—W. H. Preece.—From an English source.

Spurious Oidium Americanum in the Vineyards of France.—A biological paper. The author points out the distinctions between the *Oidium Americanum* and other vineyard pests, such as the *Erincum*.

Induction and Thermodynamics.—M. de Mertens.—A historical summary.

Chronicle of the Electric Light.—M. Ch. Vary.—The author maintains that for equal quantities of light the

Jablochkoff burner is more economical than gas by 50 to 80 per cent.

Popular Astronomy.—M. Camille Flammarion.—The nature of this letter may be understood from its title.

November 4, 1879.

Transmissibility of Rabies from Man to the Rabbit.—Human saliva is found perfectly capable of propagating this disease.

Transportation of Motive Power.—E. Venelle.—An account of the progress made towards the transmission of power by means of electricity.

The New Chloride of Calcium Battery.—M. Alf. Niaudet.—See p. 240.

Le Textile de Lyon. November 9.

M. A. Perrett gives a tabular statement of the loss sustained by silks of different growths in the removal of the "gum."

M. Jules Imbo describes the various attempts made to produce an artificial silk or to coat the fibre of China grass with a solution of silk waste. The latter operation is to a very considerable extent successful.

Reimann's Färber Zeitung,

No. 41.

A railway official is said to have died from blood-poisoning in consequence of having passed a thread of red woollen yarn through a blister on the sole of his foot. Dr. Reimann points out the improbability of the story.

No. 42.

This issue is chiefly devoted to accounts of the Arnheim and Berlin Exhibitions.

No. 43.

A piece of English calico, according to the *Wollengewerbe*, is said to be composed of:—

Cotton	53'0
Alumina	26'0
Starch	12'0
Fatty matter	2'5
Magnesium chloride	2'0
Zinc chloride	1'5
Calcium chloride	0'5
Water	2'5

Cachon de Laval is said to be capable of spontaneous ignition, which Dr. Reimann sees reason to doubt.

The cochineal harvest in the Canary Islands has suffered greatly from unseasonable rains.

No. 44.

In addition to a number of dyeing receipts, this number gives an article on the signification of patents, and accounts of the Exhibitions at Berlin and Arnheim.

Die Chemische Industrie.
No. 10, October, 1879.

Extraction of Sulphur from Sulphurous Acid and Hydrogen Sulphide.—J. Stingl and Th. Morawski.—Taken from the *Journal für Praktische Chemie*, vol. xx., p. 76.

Examination of Commercial Samples of Albumen.—Cordillot dissolves the samples in 40 parts of water, and notes the quantities of matters remaining insoluble and the consistence of the liquid. He pronounces the Russian blood albumen the best.

Researches on the Colouring Matters of Garancin.—A. Rosenstiehl.—An examination of pseudopurpurin,

purpurin, purpurin-hydrate, alizarin, garancin-orange (murexian), purpuroxanthin, and hydropurpuroxanthin.

A. E. Mézès proposes to make up pigments with a mixture of 24 parts of glue (isinglass), 534 parts glycerin, 208 water, 208 parts wax, 12½ ammonia, and 12½ resin.

Bulletin de la Société d'Encouragement pour l'Industrie Nationale.

No. 69, September, 1879.

This issue contains no chemical matter.

MISCELLANEOUS.

Royal Institution of Great Britain. — General Monthly Meeting, Monday, December 1; the Duke of Northumberland, LL.D., D.C.L., Lord Privy Seal, President, in the chair. Miss Henrietta Maria Adair, Edward Greenhill Amphlett, Esq., M.A., Henry Fearnside, Esq., M.B., F.R.C.P., Major Edward Smith Gordon, R.A., and Thomas Henry Sanderson, Esq., were elected members. The following are the arrangements for lectures before Easter, 1880:—

Christmas Lectures.

Professor Tyndall, D.C.L., F.R.S.—Six Lectures on Air and Water; on December 27 (Saturday), 30, 1879; Jan. 1, 3, 6, 8, 1880.

Prof. Edward A. Schäffer, F.R.S.—Ten Lectures on the Physiology of Muscle; on Tuesdays, Jan. 13 to March 16.

H. Heathcote Statham, Esq.—Two Lectures on Modern Architecture since the Renaissance; on Thursdays, Jan. 15 and 22.

Prof. Dewar, M.A., F.R.S.—Eight Lectures on Recent Chemical Progress; on Thursdays, Jan. 29 to March 18.

Prof. T. Rupert Jones, F.R.S.—Three Lectures on Coal; on Saturdays, Jan. 17, 24, 31.

Prof. Ernst Pauer.—Three Lectures on Handel, Sebastian Bach, and Joseph Haydn. With Musical Illustrations. On Saturdays, Feb. 7, 14, 21.

Four Lectures, on History of Literature, on Saturdays, Feb. 28, March 6, 13, 20.

The Friday Evening Meetings will begin on January 16, at 8 p.m. Prof. Dewar, F.R.S., will give a discourse (Studies on the Electric Arc) at 9 p.m. Succeeding discourses will probably be given by Dr. W. B. Carpenter, Prof. J. Marshall, Dr. Huggins, Mr. W. H. Preece, Rev. H. R. Haws, Mr. F. J. Bramwell, Mr. H. N. Moseley, Dr. C. William Siemens, Profs. Tyndall and Huxley, Lord Reay, Mr. G. J. Romanes, M. Lecoq de Boisbaudran, Mr. W. H. Pollock, Prof. Frankland, Mr. H. H. Statham, Mr. W. Spottiswoode, and Mr. Warren De la Rue. To these meetings members and their friends only are admitted.

MEETINGS FOR THE WEEK.

- MONDAY, 8th.—Medical, 8.
— Royal Geographical, 8.30.
— London Institution, 5.
— Society of Arts, 8. (Cantor Lecture). "Chemistry of Bread and Bread Making," Dr Graham.
TUESDAY, 9th.—Civil Engineers, 8.
— Photographic, 8.
WEDNESDAY, 10th.—Society of Arts, 8.
— Microscopical, 8.
THURSDAY, 11th.—Royal, 8.30.
— Royal Society Club, 6.30.
FRIDAY, 12th.—Astronomical, 8.
— Quekett Club, 8.
SATURDAY, 13th.—Physical, 3. "A New Form of Resistance Balance for Comparing Standard Coils," J. A. Fleming, D.Sc. "The Graduation of Prof. Hughes' Sonometer," J. H. Poynting. "A Dispersion Photometer," W. E. Ayrton and J. Perry. "The Value of 'g' at Tokio, Japan," W. E. Ayrton and J. Perry.

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THE CHEMICAL NEWS.

Vol. XL, No. 146.

THE PERIODIC LAW OF THE CHEMICAL ELEMENTS.

By D. MENDELÉEF.

(Continued from page 268.)

THE metallic character appears more clearly in the members of the even series, while the corresponding members in the odd series rather possess acid properties. Thus there is a marked difference between V, Nb, and Ta of the even series in the fifth group and P, As, Sb, and Bi, although they all give an oxide corresponding to R_2O_3 ; further, the first-named give acids less energetic than the latter. The members of the even series do not give, as far as is known, any hydrated or volatile metallo-organic compounds like the corresponding members in the odd series. As, among the odd elements, Zn, Cd, As, Sb, Se, Te, Br, I, Sn, Pb, Hg, and Bi can form, in a general way, metallo-organic compounds, we can foresee with certainty that the elements In and Tl, comprised amongst those which have just been cited, will also give metallo-organic compounds, such as $InAc_3$ and $TlAc_3$. Up to the present time there is no member of the even series of the higher groups which has formed a metallo-organic compound. The attempts made by Buckton, Cahours, &c., to prepare $TiAc_4$ by means of $TiCl_4$ have not succeeded, in spite of the great analogy between $TiCl_4$, $SiCl_4$, and $SnCl_4$. If, then, the even elements should give any metallo-organic compounds, these compounds would behave in an altogether different manner to the metallo-organic bodies already known, for the same reason that the hydrogenated compounds of Pd, Cu, Nb have not the same properties as the corresponding compounds of the odd series. It would be difficult to obtain volatile compounds of hydrogen and ethyl with Zr, Nb, Mo, W, and Ur.

It might seem that the establishment of the second series would be in opposition to the general division into odd and even series, because the members of this even series (Li, Be, B, C, N, O, F) possess acid properties, give hydrogenous and metallo-organic compounds (BAe_3 , $CAe_4 = C_9H_{20}$, NAe_3 , OAe_2 , FAe), and some of them are gaseous, that is to say, that they behave like the odd elements. But in any case, it is necessary to remark, relatively to this series: 1st. That it is not connected with the eighth group like the other even series are; 2nd. That the difference between the atomic weights of elements of this series and the corresponding atomic weights of the following series is about 16, while the difference for all the other series is from 24 to 28. The difference between the atomic weights of the successive even series is about 46, whilst the elements of the second and fourth series have only a difference in their atomic weights of from 32 to 36.

Li	Be	B	C	N	O	F	Na	Mg	Al	Si	P	S	Cl
K	Ca	"	Ti	V	Cr	Mn	Cu	Zn	"	"	As	Se	Br
Difference 32	31	"	36	37	36	36	40	41	"	"	44	46	45

In this manner the apparent anomalies are explained, and the fundamental idea that we have conceived concerning the dependence between the properties of the elements and their atomic weights is but confirmed by these observations. For, since there exists another difference in the atomic weights, the reciprocal relation between the properties of the elements should also be different.

The elements of the second series should not have any properties coinciding with those of the elements in the fourth series, except when they would have atomic weights less than those which they really possess. These differences are still noticed between Na and Cu, Mg and Zn; but they disappear when we compare P and As, S and Se, Cl and Br, elements whose differences of atomic weights and properties remain within ordinary limits. It is because of these differences noticed amongst the elements of the second series that I distinguish them by the name of *typical elements*. We can add Na and Mg to them, independently of H, for reasons which have been mentioned above.

If we compare the reciprocal relations of the other analogues with the relations that the compounds of the homologous series have amongst themselves, we see that the typical elements correspond to the initial members of the homologous series, which, as we know, do not possess all the properties of the higher homologues. In the same manner the initial members (H_2O and CH_4O) of the alcoholic series $C_nH_{2n} + 2O$ possess many properties peculiar to themselves, which disappear in the higher members.

From what precedes, appertains the isolated independent position of hydrogen, which has the lowest atomic weight. From the nature of the saline oxide H_2O and the salts HX , it ought to belong to the first group; the nearest analogue of hydrogen is sodium, which should be placed in an odd series of the first group. Cu, Ag, and Au are very distant analogues of H. All the five give compounds corresponding to RO and R_2O_2 (H_2O_2 , Na_2O_2 , Cu_2O_2 , Ag_2O_2 , and Au_2O_2). If oxide of copper at the minimum of oxidation is represented by CuO , it is also necessary to represent the preceding peroxides by RO , NaO , AgO , AuO . It is true that CuO forms salts corresponding to CuX_2 , while the peroxides, such as AgO , do not give, as far as is known, any salts of this kind. However, Ni, which immediately precedes Cu in the system, and Pd, which immediately precedes Ag, have an analogous difference; Ni gives no salts like NiX_2 , while Pd forms salts like PdX_2 , although they are not very stable.

Further, we notice, in each even series, from the first group to the eighth, an increase in the quantity of oxygen which can enter into combination with an element; in the series of the eighth group this faculty diminishes as the atomic weight increases (see above), and it attains its minimum for Cu, Ag, and Au; from thence (for Zn, As, for Cd, In, for Hg, Tl) it commences to increase. Therefore Cu, Ag, and Au, as is shown in the second table, occupy two places, one in the first group and the other in the eighth; in the lower forms of combination they correspond to H and Na (first group). This analogy is above all evident in Ag, which needs no other explanation; the analogy of the compounds of suboxide of copper and suboxide of gold with the compounds of oxide of silver is quite as certain; the comparison of the properties of $CuCl$, $AgCl$, and $AuCl$, afford sufficient proof. In spite of a great analogy between the compounds HX and NaX , there exists between the first and the last a series of differences well known to everybody; it calls to mind the differences between carbonic acid and the higher homologues (glycolic acids), which are analogous to it in many respects.

As a noticeable example of the analogy between the compounds of H, Na, and Cu (as suboxides), Ag and Au (also as suboxides), I will give a table of their volumes. (See next column.)

We see that the corresponding members have very nearly equal volumes. Li and K, on the contrary, have in the same form of combination different volumes. It is in this manner that the volume of $LiCl = 21$, of $KCl = 37$.

* It is known that K gives a peroxide KO_2 . It would be interesting to further the study of peroxide of lithium, to decide whether its formula is LiO , and if it has not basic properties, even in a small degree. AgO should also be studied in this respect.

† We know that I, in reacting on peroxide of sodium, forms Na_2OI , which has a composition analogous to that of Cu_2OCl .

		Density.	Volume.						
HCl	..	..	..	1.27	29				
NaCl	..	..	..	2.16	27				
CuCl	..	..	..	3.68	27				
AgCl	..	..	..	5.55	26				
AuCl	..	..	..	9.3 ?	25				
		Volume.		Volume.				Volume.	
Na ₂ CO ₃	..	43	H ₂ O	..	20	H ₂ SO ₄	..	53	
Ag ₂ CO ₃	..	46	Na ₂ O	..	21	Na ₂ SO ₄	..	54	
						Ag ₂ SO ₄	..	58	
HNO ₃	..	41	NaHO	..	19	NaClO ₃	..	46	
NaNO ₃	..	39	Cu ₂ O	..	25	AgClO ₃	..	44	
AgNO ₃	..	39	Ag ₂ O	..	28				

of LiNO₃=29, of KNO₃=48, of Li₂SO₄=50, of K₂SO₄=66. The compounds of K have greater volumes (and a constantly less density) than the corresponding compounds of Na or Li.

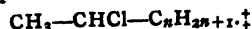
(To be continued.)

ON THE NORMAL PARAFFINS. PART III.*

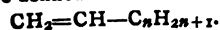
By C. SCHORLEMMER, F.R.S.,
Professor of Organic Chemistry in Owens College, Manchester.

THE isomeric monochlorides, obtained from the normal paraffins existing in petroleum, yield by the abstraction of hydrochloric acid a mixture of olefines, one portion of which readily combines with hydrochloric acid in the cold, whilst the other unites with it only on heating.†

The chlorides formed in the cold boil with partial decomposition and at a lower temperature than the others, which distil without undergoing any change, and have the general formula—



They are therefore defined from the olefines of the series—



Similar results have been obtained by Le Bel.§

The constitution of the olefines combining with the acid in the cold is not yet known. They are possibly not derived from normal paraffins, but from isomerides, which cannot be separated from the former by distillation. On the other hand, their formation can also be explained without making this assumption. This question can only be decided by using an absolutely pure paraffin.||

For this purpose normal hexane from mannite was selected, which possibly might yield three isomeric monochlorides:—

- (1.) CH₃—CH₂—CH₂—CH₂—CH₂—CH₂Cl
- (2.) CH₃—CH₂—CH₂—CH₂—CHCl—CH₃
- (3.) CH₃—CH₂—CH₂—CHCl—CH₂—CH₃.

The formation of the first two has already been proved.¶ The following seemed capable of determining whether the third is also produced. By the abstraction of hydrochloric acid three hexylenes might be formed:—

- Butyl-ethylene—
CH₃—CH₂—CH₂—CH₂—CH=CH₂
Methyl-propyl-ethylene—
CH₃—CH₂—CH₂—CH=CH—CH₃
Diethyl-ethylene—
CH₃—CH₂—CH=CH—CH₂—CH₃.

The first of these does not combine with cold hydrochloric acid; the second is the hexylene obtained from secondary hexyl iodide.¶ Le Bel and Wassermann have

* Abstract of a Paper read before the Royal Society, November 20, 1879.

† Journ. Chem. Soc., 1873, p. 319.

‡ Morgan, *ibid.*, 1875, p. 301.

§ Bull. Soc. Chim. (2), xxviii., p. 460.

¶ Journ. Chem. Soc., 1875, p. 306.

¶ Hecht, *Dent. Chem. Ges. Ber.*, xi., p. 1152.

found that cold hydrochloric acid has no action on it, from which it follows that, if normal hexane from mannite yields a hexylene combining with hydrochloric acid in the cold, it could only be diethyl-hexylene, which could be easily identified by conversion into ethyl-propyl carbinol and oxidising it, when only propionic acid should be formed.

This was my programme; the results were, however, quite unexpected.

The hexylene obtained by decomposing the hexyl chlorides was left in contact with cold fuming hydrochloric acid for some weeks. *The whole of it combined, and the hexyl chloride thus formed boiled constantly and without the least decomposition at 124° to 125°.* It was converted into the alcohol, which on oxidation yielded only acetic acid and butyric acid, and consequently is methyl butyl carbinol. We have therefore the remarkable fact that two hexanes exist, which must be regarded as normal compounds, and therefore, according to our present theory, to be identical. This is, however, not the case. I have already, in my first paper, pointed out some other differences existing between the two hexanes, but left the question open whether these are caused by impurities contained in the hexane from petroleum, or whether we have here a case of fine isomerism, for which an explanation has to be found.* I believe the results of my present research speak strongly in favour of the latter view.

For several reasons I am inclined to believe that petroleum consists chiefly of an inextricable mixture of isomeric and homologous paraffins, in which, however, the normal hydrocarbons preponderate. This would certainly explain why it is so difficult to isolate from it bodies having a constant boiling-point,† but not the differences exhibited by the two hexanes.

A combination of these researches has already been commenced. My friend Thorpe, who has made the most interesting discovery that the terebinthinate exudation of *Pinus Sabiniensis* contains a large quantity of normal heptane,‡ has kindly offered me to join him in the chemical investigation of this hydrocarbon. At the same time we shall compare it with other "normal" heptanes from different sources.

ON THE ACTION OF NUCLEI IN PRODUCING THE SUDDEN SOLIDIFICATION OF SUPERSATURATED SOLUTIONS OF GLAUBER'S SALT.§

By CHARLES TOMLINSON, F.R.S.

In this paper the author states the general conclusions at which he has arrived since resuming the study of this subject two years ago.|| The remarks that are made refer to solutions of the typical salt, sodic sulphate, in the proportions of 6 of salt to 3 of water.

In his first two papers on supersaturated saline solutions¶ a nucleus is defined as a body that has a stronger adhesion for the salt, or for the liquid of a solution, than subsists between the salt and the liquid. Many examples are given in which oils, fixed and volatile, alcohol, ether, &c., spreading on the surface of the solution, lead to the separation of salt and the solidification of the solution; while if such liquids, instead of spreading, assume a lenticular form, they become inactive, and by shaking the flask with a circular motion are dispersed through the solution in numerous globules, without any immediate nuclear action.

* Phil. Trans., vol. clxii., p. 119.

† Journ. Chem. Soc., 1875, p. 306.

‡ *Ibid.*, 1879, p. 236.

§ Abstract of Paper read before the Royal Society, November 20, 1879.

¶ Proc. Roy. Soc., vol. xxvi., p. 523.

¶ Phil. Trans., 1868 and 1871.

There are several circumstances which favour the action of these liquid nuclei, such as (1) chemically clean flasks and solutions, so as to maintain (2) the surface-tension of the solution as high as possible, in order to spread the oils; (3) bright and clear weather, with strong evaporative force. Under such conditions the oils, &c., usually form films with nuclear action.

But it has been shown that in closed flasks, with special precautions against the entrance of particles from the air, the oils, &c., may be made to spread on the surface of these solutions without any nuclear effect.

The author was led to make a number of observations daily, during some months, on the action of a freshly-distilled volatile oil on solutions of sodic sulphate, and found that during certain winds the oil was active, and that while other winds prevailed it was passive.*

It was clear, then, that during some winds a force was present in the air which rendered the oil active; while during other winds, the force being absent, the oil remained passive. He succeeded in identifying this force with ozone; and the point now to be determined is, what is the peculiar function of ozone in rendering oils, &c., active, as nuclei?

It is evident, from the behaviour of ozone on volatile oils exposed to its influence, that its action is an oxidising one, diminishing their cohesion, just as rust weakens the cohesive force of the particles in a bar of iron.

The effect which ozone produces quickly on a volatile oil is produced slowly on the same oil by long keeping. Its cohesive force is so far weakened, if not destroyed, that a drop of it on the surface of a solution no longer tends to assume the lenticular form, or to disperse in globules through the solution. Under such circumstances, since oil adheres much more strongly to salt than to water, and the supersaturated solution, being a highly charged system, capable of yielding to a force that is exerted in the right direction,† such an oil is capable of adhering to and separating a portion of the salt, and the action, once begun, is propagated rapidly, until the whole solution is solid.

The action of a foreign nucleus introduced into the cold solution from without, is to determine the formation of the ten-atom or normal salt; but if a solid body be boiled up with the solution, and the covered flask be left to cool to temperatures about and below 40° F., the modified or seven-watered salt will be found clustered about the solid body. An explanation of this may be found in the behaviour of the salt while undergoing solution. Suppose the flask to contain 6 parts of the crystallised, not effloresced, salt and 3 of water. If the heat be applied too strongly the salt gives up its water of crystallisation, and the anhydrous salt is thrown down, and produces violent bumpings of the flask. But if the heat be kept below the temperature of maximum solubility of the salt (about 93° F.) until the whole of it is dissolved, the solution may be raised to the boiling-point and also be cooled down in covered vessels to about 40° F. without any separation of salt. The anhydrous salt now appears to be in solution,‡ and a solid body previously heated in the solution forms a portion of it, or rather a portion of the flask, and may be regarded as a prolongation of its sides. Under a proper reduction of temperature, crystals of the seven-atom salt will cluster about such a body, but it cannot be said to exert any active nuclear function. It was pointed out by Ziz, as long ago as the year 1809, that solid bodies boiled up with the solution are inactive; and

the author has found them to be so when put into the hot or warm solution, and left till cold; and that oils and other liquids, boiled or heated with the solution, become equally inactive. Several observers, who have more or less recently worked on the subject before us, have not attended to this condition, and hence, obtaining negative results, have declared that the oils, &c., have no nuclear action under any circumstances.

If, however, a nucleus from without be introduced into the cold solution, a nucleus that is capable of adhering to the salt and not to the water of the solution, a molecule of the anhydrous salt is thereby set free, and, in the very act of separating, 10 atoms of water enter into combination, and thus determine the solidification of the whole of the solution in the form of the normal salt.*

If the nucleus fail to catch one of these saline molecules, as it were, in the nascent state, but disperses through the solution, its effect is to lessen the adhesion between the water and the salt, and a portion of the seven-atom salt is liberated (mostly at the bottom of the vessel, where the solution is richer in salt) provided the temperature be not too high: this effect is also produced by cold alone.

If an essential oil, which acts as a nucleus in determining the formation of the normal salt, be submitted to distillation, the distillate re-assumes its cohesive force, and when added to the solution is no longer active, seeing that whether in the lenticular form or dispersed in globules it has a separate existence of its own, the lens and each minute globule being bound up in their own surface tension, which prevents them from coming into contact with the salt. If, however, the drop of oil spread out into a film upon the surface of the solution, the cohesion is so far weakened that adhesion between it and the salt becomes possible, separation usually takes place, and large crystals of the normal salt mould themselves on the under surface of the film.

If such be the true explanation, it is clear that any other force which diminishes or destroys the cohesive force of the oil must confer upon it the same nuclear property. Consequently a freshly distilled essential oil, mineral naphtha, &c., which are inactive, become active by the addition of a little absolute alcohol, ether, and similar liquids in which they are soluble.

The fixed oils which tend to become rancid through age are also active, and the stearine deposited by them is especially so. Newly refined fixed oils, which are inactive, become active on the addition of a little alkali; but the tendency of potash, soda, and ammonia is to form soluble soaps with the oils, which mingle with the solution often without nuclear action. Lime and magnesia are more manageable. On mixing the oil with one of these, and adding a drop or two to the solution, then inclining the flask nearly horizontally, and rotating it slowly on its axis, these insoluble soaps adhere to the side and form smears, upon which masses of large crystals of the normal salt start into existence in a striking manner, and the solution immediately becomes solid, so that on restoring the flask to its erect position a vertical wall appears, with a solid mass on one side and an apparently empty space on the other.

Some of the solid and semi-solid fats in their natural state, or as they leave the hands of the cook, are admirable nuclei. Such are suet, dripping, fat of bacon and ham, lard, butter, and some others. But in all cases a freshly cut surface acts most effectually, and a fat that sometimes does not act for hours if cut one way, will act instantly if cut at right angles to the former direction. Lard contains a good deal of entangled water: this should be driven off. The lard when cold sometimes acts better when smeared on a clean glass rod than in a small lump.

The action of solid porous nuclei, such as plaster of

* Löwell noticed that when the transparent seven-atom salt is removed from the solution, it becomes opaque and hot in consequence of the fixation of three additional atoms of water. In such case the supersaturated mother-liquor on the surface, and entangled among the crystals, produces, in the absence of a nucleus, a crystal of the normal salt by evaporation, and this acts as a nucleus.

* These observations were made at Highgate; others by the sea-side are described in *Proc. Roy. Soc.*, No. 196.

† See a Paper "On the Function of the Sides of the Vessel in maintaining the State of Supersaturation." *Proc. Roy. Soc.*, vol. xxvii., p. 189.

‡ See *CHEMICAL NEWS* (vol. xx., pp. 267 and 277). In Wüllner's experiments on the elasticity of steam, when sodic sulphate was added to the water, the diminution in the elasticity was found to be proportional to the quantity of dry salt in solution at temperatures from 10° to 100° C. At the point of maximum solubility of the salt no molecular change occurred, or it would have impressed itself on the curve which represented the elasticity of the steam.

Paris, pumice, &c., in determining the sudden solidification of these solutions admits of easy explanation. When these bodies are moderately absorptive they act by separating water from the solution; but when, by exposure to heat, their absorptive power is considerably increased, they become immediately saturated, not with water, but with a portion of the solution itself, and hence there is no separation of salt. When such an apparently inactive body is exposed to the open air, its absorptive power becomes somewhat diminished by access of moisture, and it is brought into a condition to absorb more water from the solution, instead of the solution itself. In the experiments of Mr. Liversidge and others the porous bodies were thoroughly heated and dried before they were brought into contact with the solution, and this will sufficiently explain the negative results obtained by them in their carefully conducted experiments.

The action of nuclei on the supersaturated solutions of other salts, such as those of alum, zincic and magnesian sulphates, and one or two others, admits of similar explanations; but there are peculiarities belonging to each which need not at present occupy the time of the Society.

The curious condition of supersaturation was first observed by Dr. Black towards the end of the last century, and from that time to the present innumerable notes and memoirs have been written on the subject. Some observers who have treated the subject most elaborately have adopted methods which, by the action of heat and the exclusion of air, have tended to promote the cohesive or adhesive force of the bodies examined by them, and obtaining only negative results, and always a certain result with a crystal of the salt, they have insisted that this is the only nucleus. Others, again, have sought for an explanation in some catalytic or other mysterious force; while a third set of observers have declared it to be a matter of uncertainty or hazard whether a foreign body acts as a nucleus or not. In reviewing the subject and repeating his experiments in various ways, the author sees no reason for withdrawing from the theory which he submitted to the Royal Society eleven years ago, namely, that the action of nuclei is simply mechanical, and is capable of being expressed by the familiar word "adhesion."

THE ACCUMULATION OF NITRIC NITROGEN IN THE UNCHANGED WATER OF AQUARIA.

By H. WILLIAMS JONES, F.C.S.

IN many public aquaria, as at the Crystal Palace and Westminster, and also at the very fine one recently opened at the Aston Lower Grounds, near Birmingham, unchanged sea-water is employed; and such water has been found by experience to be fit for an almost indefinite period for maintaining marine animals in health. The principle adopted is to keep the bulk of the water in underground reservoirs, and to constantly pump from them a fresh supply into the show tanks, the delivery of the water being made through a series of fine jets. The great object of a constant supply, delivered in such a manner, being to force air along with fresh cool water into the tanks containing the animals. By that means the water is maintained of an almost constant temperature, and even if temporarily rendered cloudy by the presence of decaying fragments of food, &c., the rapid oxidation resulting from the air forced in along with the fresh water from the jets, soon renders the water quite clear. At Brighton and some other places the sea-water is intended to be used for a limited time only, a fresh supply being pumped in from the sea as often as required. Since it would be impracticable for inland aquaria to be supplied even at distant intervals with fresh sea-water, on account

of the great cost incurred in collecting and for carriage, the system of unchanged water, has of necessity to be adopted. It occurred to me to examine a sample of water in my possession, which, after being used for eight years in a large public aquarium, was quite bright and appeared to answer perfectly the requirements of the animals, to see how far the nitric nitrogen had accumulated. It was inferred that a very large amount would be present as a result of the oxidation of the organic matter thrown off by the animals, or left in the water in the form of uneaten and unremoved dead food. As anticipated, a very large excess over what exists in actual sea-water fresh from the ocean was detected and the following figures will show the difference between the aquarium-water and sea-water taken about the same date from Brighton, and which fairly represents the amount present naturally in sea-water.

		Nitric Nitrogen. Parts per 100,000.	
Recent sea-water		0.0325	
Aquarium-water		12.0498	

In all well-stocked aquaria the number of living animals of a large size in relation to the bulk of water is greatly in excess of what exists in Nature; and the food of such animals kept in confinement instead of being produced in the water in which they live is received from an outside source in the shape of food formed in the ocean. The nitric nitrogen is formed in aquaria faster than it is removed by vegetation, which is usually limited to *conferve*, and so it accumulates.

ON THE VAPOUR-DENSITIES OF PEROXIDE OF NITROGEN, FORMIC ACID, ACETIC ACID, AND PERCHLORIDE OF PHOSPHORUS.

By J. WILLARD GIBBS.

(Continued from p. 236.)

Acetic Acid.—For this substance the densities have been calculated by the formula—

$$\log. \frac{2.073 (D - 2.073)}{(4.146 - D)^2} = \frac{3520}{t_c + 273} + \log p - 11.349, \quad (12)$$

the constants 3520 and 11.349 being derived from the determinations of Cahours and Bineau, which with those of Horstmann and Troost are given in Table IV. The experiments of Cahours and Horstmann were made under atmospheric pressure, those of Horstmann* by the method of Bunsen, those of Cahours presumably by the method of Dumas. The numbers in the first column of the densities observed by Cahours are taken from the twentieth volume (1845) of the *Comptes Rendus*, except a few cases distinguished by parentheses, which are taken from the preceding volume (1844). The numbers in the second column are taken from his "Leçons de Chimie générale élémentaire," 1856. These numbers seem to be based in part upon new experiments, and in part upon a revision of the observations recorded in the *Comptes Rendus*, the calculations being carried out to another figure of decimals. They are therefore entitled to a greater weight than the numbers of the preceding column.

The agreement of the formula with the numbers given in the "Leçons de Chimie" is very good, the greatest divergences being 0.080 at 190° and 0.062 at 180°. But at 190° the table in the *Comptes Rendus* agrees precisely with the formula, and at 171° (the next experiment) it shows a divergence in the opposite direction. The next divergencies in the order of magnitude are -0.033, -0.036, -0.032 at 219°, 231°, 240° respectively. Here the table in

* *Leçons de Chimie*, suppl. vi., p. 65.

TABLE IV.—ACETIC ACID.

Experiments of Cahours, Horstmann, Bineau, Troost.

Tempera- ture.	Pres- sure.	Density calc. by eq. (12).	Density observed.			Excess of observed Density.		
			Cahours.		Horst- mann.	Cahours.		Horst- mann.
			C. R.	Lecons.		C. R.	Lecons.	
338	(760)	2'077	2'08			'00		
336	(760)	2'077		2'082			+ '005	
327	(760)	2'078	2'08	2'085		'00	+ '007	
321	(760)	2'079	2'08	2'083		'00	+ '004	
308	(760)	2'081		2'085			+ '004	
300	(760)	2'082	2'08			'00		
295	(760)	2'084		2'083			- '001	
280	(760)	2'089	2'08			- '01		
272	(760)	2'093		2'088			- '005	
254'6	747'2	2'105			2'135			+ '030
252	(760)	2'108		2'090			- '018	
250	(760)	2'111	2'08			- '03		
240	(760)	2'122		2'090			- '032	
233'5	752'8	2'132			2'195			+ '063
231	(760)	2'137	(2'12)	2'101		(- '02)	- '036	
230	(760)	2'139	2'09			- '05		
219	(760)	2'165	2'17	2'132		+ '01	- '033	
200	(760)	2'239	2'22	2'248		- '02	+ '009	
190	(760)	2'298	2'30	2'378		'00	+ '080	
181'7	749'7	2'359			2'419			+ '063
180	(760)	2'376		2'438			+ '062	
171	(760)	2'466	2'42			- '05		
170	(760)	2'477		2'480			+ '003	
165'0	754'1	2'534			2'647			+ '113
162	(760)	2'575		2'583			+ '008	
160'3	751'6	2'594			2'649			+ '055
160	(760)	2'601	2'48			- '12		
152	(760)	2'716	(2'72)	2'727		('00	+ '011	
150	(760)	2'747	2'75			'00		
145	(760)	2'826	(2'75)			(- '08)		
140	(760)	2'910	2'90	2'907		- '01	- '003	
134'3	748'8	3'001			3'108			+ '107
131'3	754'1	3'055			3'070			+ '015
130	(760)	3'082	3'12	3'105		+ '04	+ '023	
128'6	752'9	3'103			3'079			- '024
125	(760)	3'168	3'20			+ '03		
124	(760)	3'185		3'194			+ '009	
			Bineau. (2'86)		Troost.	Bineau. (- '19)		Troost.
132	757	3'05						
130	59'7	2'31			2'12			- '19
130	30'6	2'21			2'10			- '11
129	633	3'03	(2'88)			(- '15)		
36'5	11'32	3'03		3'62		- '01		
35'0	11'19	3'65		3'64		- '01		
30'0	6'03	3'61		3'60		- '01		
28'0	10'03	3'75		3'75		'00		
24'0	5'75	3'71		3'70		- '01		
22'0	8'64	3'82		3'85		+ '03		
22	2'70	3'59		3'56		- '03		
21'0	4'06	3'70		3'72		+ '02		
20'5	10'03	3'86		3'95		+ '09		
20'0	8'55	3'84		3'88		+ '04		
20'0	5'56	3'77		3'77		'00		
19'0	4'00	3'73		3'75		+ '02		
19	2'60	3'65		3'66		+ '01		
12'0	5'23	3'88		3'92		+ '04		
12	2'44	3'77		3'80		+ '03		
11'5	3'76	3'84		3'88		+ '04		

the *Comptes Rendus* agrees substantially with that of the "Leçons," but the experiments of Horstmann show a divergence to the opposite direction. In fact, the three columns of observed densities nowhere agree in the direction of their divergence from the formula.

The somewhat decided differences between the results of Horstmann and those of Cahours may be due in part to the different methods of observation; especially to the entirely different manner of applying the heat and measuring the temperature, but the higher values obtained

by Horstmann cannot be accounted for by too short an exposure to the source of heat, for his experiments were made with decreasing temperatures.

The determinations of Bineau are taken from the same sources as those on formic acid, the earlier determinations being distinguished as before by parentheses. One of these (at 132°) was made by the method of Dumas, the other by that of Gay-Lussac. The smallness of the observed densities appears due to the presence of water (An acidimetric test gave 39 parts of acid in 100.) The

TABLE V.—ACETIC ACID.

Experiments of Naumann.

		TEMPERATURE.								
		78°.	100°.	110°.	120°.	130°.	140°.	150°.	160°.	185°.
A	Pressure		393.5	411	432	455	477	498.5		565
	Density calculated		3.39	3.23	3.06	2.90	2.75	2.61		2.28
	Density observed		3.44	3.31	3.14	2.97	2.82	2.68		2.36
	Excess of density observed..		+0.05	+0.08	+0.08	+0.07	+0.07	+0.07		+0.08
B	Pressure		342.3	359.3	377.5	398.5	417.5	436.5		495
	Density calculated		3.35	3.18	3.02	2.85	2.70	2.57		2.26
	Density observed		3.37	3.22	3.06	2.89	2.75	2.63		2.31
	Excess of density observed..		+0.02	+0.04	+0.04	+0.04	+0.05	+0.06		+0.05
C	Pressure		258							382
	Density calculated		3.26							2.22
	Density observed		3.17							2.25
	Excess of density observed..		-0.09							+0.03
D	Pressure		232		252	274	287.5	300		335
	Density calculated		3.23		2.87	2.72	2.58	2.46		2.21
	Density observed		3.12		2.94	2.68	2.54	2.44		2.23
	Excess of density observed..		-0.11		+0.07	-0.04	-0.04	-0.02		+0.02
E	Pressure	164	186	197	209	221	232	243	253	269
	Density calculated	3.53	3.15	2.97	2.81	2.65	2.52	2.41	2.32	2.18
	Density observed	3.41	3.06	2.91	2.75	2.61	2.50	2.40	2.31	2.22
	Excess of density observed..	-0.12	-0.09	-0.06	-0.06	-0.04	-0.02	-0.01	-0.01	+0.04
F	Pressure	149	168			201				
	Density calculated.. . . .	3.50	3.12			2.62				
	Density observed	3.34	3.01			2.56				
	Excess of density observed..	-0.16	-0.11			-0.06				
G	Pressure	137	156	166.5	180	188	199	208.2		230
	Density calculated	3.48	3.09	2.92	2.75	2.60	2.47	2.37		2.17
	Density observed	3.26	2.98	2.81	2.61	2.50	2.40	2.29		2.14
	Excess of density observed..	-0.22	-0.11	-0.11	-0.14	-0.10	-0.07	-0.08		-0.03
H	Pressure	113	130	138.5	149	157.5	168.2	175		191.5
	Density calculated	3.42	3.03	2.85	2.69	2.55	2.43	2.33		2.15
	Density observed	3.25	2.94	2.78	2.60	2.47	2.32	2.26		2.13
	Excess of density observed..	-0.17	-0.09	-0.07	-0.09	-0.08	-0.11	-0.07		-0.02
J	Pressure	80	92	98.5	106	112.5	117.3		129.2	
	Density calculated	3.32	2.91	2.73	2.58	2.45	2.35		2.21	
	Density observed	3.06	2.76	2.61	2.46	2.34	2.27		2.11	
	Excess of density observed..	-0.26	-0.15	-0.12	-0.12	-0.11	-0.08		-0.10	
K	Pressure	66	77.7	84	89.5	93	98	103		110.5
	Density calculated	3.26	2.85	2.68	2.53	2.40	2.31	2.24		2.12
	Density observed	3.04	2.66	2.49	2.37	2.32	2.24	2.16		2.11
	Excess of density observed..	-0.22	-0.19	-0.19	-0.16	-0.08	-0.07	-0.08		-0.01

other experiments were made with the same apparatus which was used with formic acid, and show even greater regularity in their results than the experiments with that substance. Only in one case is the influence of proximity to saturation seen, viz., at 20.5° and 10.03 m.m., the pressure of saturated vapour at this temperature being about 12.7 m.m.* In the remaining fifteen observations of this series, notwithstanding the very low pressures employed from 2.44 to 11.32, the greatest difference between the observations and the formula is 0.04, and the average difference 0.02.

The two observations by Troost† were made by the method of Dumas, but at pressures very low for this method. The results obtained differ considerably from the formula, but not so much as in the case of his experiments at low pressure with peroxide of nitrogen.

Table V. contains the experiments of Naumann‡ on acetic acid. These consist of ten series (distinguished by the letters A, B, C., &c.) of observations by Hoffmann's method.|| The temperatures of the observations in the

different series are for the most part the same, so that for each temperature we have observations through a wide range of pressures. Within each compartment of the table are given in order the pressure of an experiment, the density calculated by equation (12), the observed density, and the excess of observed density, the temperature of the experiment being given at the head of the column. These experiments, taken by themselves, seem to show an effect of pressure upon the density about one-third greater than is indicated by the formula. But the divergences (of which the greatest is 0.26 and the average 0.085) are not large in view of the fact that the experiments were undertaken rather with the desire of obtaining a great number of observations with moderate labour than with the intention of attaining the greatest possible accuracy.

The quantity of acid diminishes somewhat regularly from 0.2084 grm. in series A to 0.0185 in series K. The volume, which was 154 c.c. in the experiment at 185° in series A, diminishes in the successive series, and in the same series with diminishing temperature to 69.6 c.c. in the experiment at 78° in series K. It is worthy of notice that the greatest deviations from the formula occur where the liability to error is most serious with respect to pressure (which was measured without a cathetometer), to volume, and to the quantity of acid.

* This number is obtained from data given by Bineau by the same kind of interpolation which was used for formic acid.

† *Comptes Rendus*, vol. lxxvi. (1878), p. 1395.

‡ *Leipzig's Annalen*, vol. clv., p. 345.

|| This is a modification of the method of Gay-Lussac, in which the heat is supplied by a vapour-bath.

Far more serious than the absolute amount of these divergences is the regularity which they exhibit. But it must be remembered that the observations are by no means entirely independent, and many sources of possible error, such as the calibration of the tube and the determination of the quantity of acid, might affect the results with considerable regularity.

Only to a slight degree can the divergences from the formula be accounted for by an insufficient exposure to the temperature of the experiment. The observations, except those at 78°, were made with increasing temperatures, and the greatest divergences from the formula are not in the positive direction. Yet the positive divergences occur where we should most expect to find them if they were due to this cause, viz., in the series in which the greatest quantities of acid were used, and in cases in which the temperature seems to have been raised at once an usual number of degrees. (See especially the observation at 120° in series D, and in general the observations at 185°, which exhibit, if not a positive, at least a diminution of negative excess.) In the observations at 78° which were the last of each series, and therefore followed a fall of temperature from 185°, we find in some cases, especially in series G, H, and J, a negative divergence much greater than in the other determinations of the same series, and which appears to be referable to this circumstance.

(To be continued.)

SPECTROSCOPIC INVESTIGATIONS.

As previous researches of G. L. Ciamician have shown, chemically related elements have homologous spectra; that is, the individual spectra of the elements of any given group differ from one another only in having their lines displaced towards one end or the other of the spectrum. In the course of a comparative investigation of the spectra of the metals of the alkaline earths, Ciamician has observed phenomena which are adapted to furnishing an explanation of these remarkable relations between the spectra of related elements. When the spectra of the metals of the alkaline earths are produced by allowing the induction spark to pass between the metals (as electrodes) with inserted Leyden jar in an atmosphere of hydrogen, spectra are obtained which show the homology of the lines very beautifully. The spectrum of magnesium, however, can not be compared with the spectra thus obtained, because it does not contain the less refrangible lines. If the jar is removed, or if a weaker battery and a smaller induction coil is used, all the lines in the red and yellow field in the spectra of calcium and strontium become less intense, and spectra are obtained which are very similar to that of magnesium. If the less refrangible part of the spectrum of the group of metals of the alkaline earths—that part which is visible only at a high temperature, corresponding to a high electric tension—is compared with the less refrangible half of the complete oxygen spectrum, it is noticed that these two half-spectra exhibit a remarkable similarity (homology). From this it would follow that the spectrum of the group of the metals of the alkaline earths appear to be made up of that of magnesium, and of the less refrangible half of the oxygen spectrum. As it is known that the atomic weights of barium, strontium, and calcium can be made up of the atomic weight of magnesium and of oxygen ($24 + 16 = 40$ [calcium], $24 + 4 \times 16 = 88$ [strontium], and $24 + 7 \times 16 = 136$ [barium]), experiments were undertaken to discover whether these remarkable relations in the spectra have any real significance. The author has shown that there exists the same kind of relation between the spectra of carbon and of nitrogen and that of cyanogen, and between the spectra of carbon and of oxygen and the spectrum of carbon monoxide, as that observed between the spectra of magnesium and of oxygen and that of the group of the metals of the alkaline earths. It can hence be said that in all probability

the homology of the spectrum lines of chemically related elements depends upon the fact that, in accordance with the regularities in the atomic weights noticed by Mendeleeff, the elements of such natural groups consist of the same components.—(*Wiener Sitzungsberichte*, 79, II. *Abth.*)—*American Chemical Journal*.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, December 4, 1879.

Mr. WARREN DE LA RUE, President, in the Chair.

THE following certificates were read for the first time:—J. C. Evans, W. H. Glazier, K. W. Hedges, R. Howell, J. Hogarth, J. McCarthy, A. K. Miller, W. O. Prosser. An election of Fellows was to have been held, but as less than forty Fellows were present the ballot could not be taken.

THE PRESIDENT expressed a hope that all members would endeavour to be present at the next meeting (Dec. 18), when, if possible, the ballot would take place.

The following papers were read:—

"*The Comparative Value of Different Methods of Fractional Distillation*," by F. D. BROWN. When a fractional distillation is carried out on a large scale either or both of two well-defined processes can be used. In the first, "washing," the mixed vapours are passed through several layers of liquid obtained by their own partial condensation; in the second, "cooling," the mixed vapours are partially condensed by allowing radiation to take place or by passing them through a coil kept at a given temperature. In both processes the liquids of highest boiling point are kept back, and a better distillate is accordingly obtained. A possible explanation of the success of the first process is that by obstructing the passage of the vapour the successive layers of liquid give it more time to cool. From this point of view the two processes are identical. In the present paper the author has endeavoured to determine whether there is any essential difference between "washing" and "cooling" or not. The mixture employed throughout these experiments was the one used by the author in his previous research, and consisted of carbon disulphide and benzene. The dephlegmator advocated by Linnemann, Bel. and Henninger, &c., was first compared with a glass tube 690 m.m. long, inclining gently upwards, exposed freely to the cooling influence of the atmosphere, &c. As a result of several experiments it was found that the distillation with the long cooling tube yielded slightly better results than that in which the vapour was washed by passing through the dephlegmator, and that both the dephlegmator and the cooling tube produced better distillates than the retort or the flask with T-piece usually employed. These results were to some extent, however, vitiated by the fact that the distillations were not carried out in equal times, and in order to render the results more strictly comparable the author constructed a long glass tube, and fitted it up with a movable series of disks of wire gauze (each disk containing a short brass tube, trapped at the bottom to prevent the vapour passing up the tube instead of through the layer of liquid condensed on the wire gauze), so that this tube could be used either as a simple cooling-tube, or by inserting the disks could be converted into a dephlegmator. Great care was taken to make the times of the distillation as equal as possible. As a result of these experiments the author concludes that "washing" and "cooling" are not identical processes, that the dephlegmator has a special value, and gives better results, other things being equal, than the simple cooling-tube. The author then

contrived a more exact and delicate process of cooling, by means of which the still-head could be kept at the lowest possible temperatures compatible with the passage of vapour into the condenser. This was effected by passing the worm of the still through a strong copper box, which is partly filled with a liquid boiling about the temperature desired. The vapour from this boiling liquid is condensed in a worm which returns the liquid to the box, the box, worm, &c., being air-tight, and constructed of stout copper. The final adjustment of temperature was obtained by connecting this air-tight system with a pump, so that the pressure under which the liquid boiled in the box could be varied at will. By this means the vapour round the still-head could be kept at a constant temperature for an indefinite period, and could be varied without much trouble by alteration of the pressure. By using this apparatus, and charging the box with carbon disulphide, and the still with a mixture containing 42.69 per cent of CS_2 , a distillate was obtained containing in two experiments 97 to 99 per cent of CS_2 . The author also tested the apparatus by distilling commercial benzene and separating almost pure benzene. Thus 1100 c.c. of a very impure sample gave (the still-head being kept at 81°) at 80.4° , 155 c.c.; 80.8° , 176 c.c.; 81° , 180 c.c.; 82° , 191 c.c. After 500 c.c. had distilled over the experiment was stopped. A sample of the residue distilled as follows:—At 95° , 89 c.c.; 100° , 130 c.c.; 105° , 158 c.c.; 110° , 186 c.c. The paper contains tables of the details of the experiments, curves showing the compositions of the distillates, and drawings of the apparatus.

Dr. ARMSTRONG said that as far as he knew no one had previously compared the values of these two processes of washing and cooling from a scientific point of view. He could confirm Mr. Brown's statements as to the importance of the rate of distillation and the extent of cooling surface. He did not think that fractional distillations of homologous bodies, the vapours of which seemed, so to speak, to hang together, were of much value unless large quantities were employed.

Mr. NEILSON thought that a separation could be effected by fractionally distilling small quantities. He had used a very simple apparatus with good results. It consisted of a flask, the neck being closed with a cork, through which passed a tube ending inside the flask in a bulb, with a drawn-out and turned-up beak.

The CHAIRMAN remarked that he had worked on the same principle as that carried out so completely by the author; but instead of surrounding the still-head with a vapour of constant temperature, a thermometer was inserted in the still-head, and the temperature carefully watched.

Mr. BROWN did not think that any hanging together of vapours took place. In his opinion, small quantities could be successfully manipulated by using an apparatus of small size.

The next paper was read by Mr. MUIR, "On the Influence Exerted upon the Course of certain Chemical Changes by Variations in the Amount of Water of Dilution," by M. M. P. MUIR and C. SLATER. The authors have studied the influence of the addition of water on the reactions which ensue when solutions of (1) calcium chloride and sodium carbonate, (2) strontium chloride and sulphuric acid, and (3) barium chloride and potassium oxalate are respectively mixed at various temperatures and for various times. The two solutions under experiment were mixed, allowed to stand for the required time, and the precipitate formed estimated by filtering or by pipetting and titrating some of the clear supernatant liquid. In the first reaction the amount of chemical change decreases regularly as the dilution increases; in the second and third various irregularities occur, which the authors have investigated in detail. The paper is lengthy, contains numerous and elaborate tables and curves, without which the results cannot be given intelligibly. The authors continue—Every chemical system appears to tend towards that condition of equilibrium which is marked by the greatest loss of energy. This

tendency may be arrested in various ways, e.g., by impressing upon the system what may perhaps be described as an artificial state of equilibrium. Thus, the condition of most stable equilibrium for a system originally consisting of barium chloride and potassium oxalate would be that in which barium oxalate and potassium chloride are produced; but by adding much water a portion of the reacting molecules is, according to the hypothesis of the authors, loaded with water of hydration. In this loading energy is lost, but the loss is less than that which would occur were molecules of barium oxalate and potassium chloride formed. The system is therefore unstable, but a certain degree of stability is impressed upon it by the presence of a large mass of one of the products of the dissociation of the complex and unstable hydrated molecules. Thus a system in a state of strain is produced: a small force may be sufficient to relieve the strain, and the relief may be attended with a rapid re-arrangement of the parts of the system. Accordingly the authors found that small physical differences, such as the roughness or smoothness of a beaker, filtering instead of pipetting, &c., had a large influence in dilute solutions. The authors conclude with the hypothesis that the amount of chemical change which occurs when barium chloride and potassium oxalate are mixed in the proportion of 1 : 1 molecule is irregularly affected by the mass of water of dilution present, because the entire system is brought into a state of strain due to the stress between its parts, and that the principal forces of which this stress is compounded are,—the force tending to produce cryohydrates and other hydrated molecules, the force tending to split up these molecules, and the force tending to separate and so to impart greater mobility to the chemically active molecules of the system.

Mr. MUIR then read a paper "On the Influence of Temperature upon the Decomposition of Barium Chloride by Potassium Oxalate in Aqueous Solutions." In concentrated solutions temperature has but little influence; with more dilute solutions the first increase of about 20° causes a marked increase of action, after this the influence of temperature is more regular; with still more dilute solutions increase of temperature causes at first a slight increase, then a rapid increase of action, until a point is reached, after which further increase of temperature but slightly affects the amount of chemical change; with very dilute solutions the influence of temperature again becomes nearly regular.

Mr. NEILSON showed, by the aid of some mathematical formulæ, that for the lower temperatures a longer period of time should be given, otherwise the results obtained would not be comparable with those obtained at the higher temperatures.

The next paper was read by Dr. F. R. JAPP, "On α - and β -Phenanthrene Carbonic Acids." The author has already described with Dr. Schultz (*Deuts. Chem. Gesell.*, x., 1861) the α acid; he has since prepared it in a purer state, and finds its melting-point to be 266° instead of 260° . This acid was prepared from crystallised calcic phenanthrene sulphonate by converting the latter into the sodium salt, distilling the dry sodium salt with potassic ferrocyanide, and saponifying the nitrite thus obtained. In the preparation of the calcic phenanthrene sulphonate much dark coloured syrupy mother-liquor was left. This mother-liquor was subjected to the same processes, which had yielded α -phenanthrene carbonic acid from the crystallised salt, in the hope that the corresponding phenanthrene carbonic acid might be more easily purified than the syrupy sulphonate; 80 grms. of crude acid were thus obtained from 2 kilos. of commercial phenanthrene. After considerable difficulty the acid was purified by recrystallising it as a sodium salt. By distillation with soda lime phenanthrene was obtained, being identified by its melting-point, picric acid double compound, and the quinone obtained by oxidation. By oxidation with chromic anhydride in acetic acid phenanthrene quinone was formed. The author has tabulated a comparison of the salts of the α and β acids. The α acid crystallises from hot glacial

acetic acid in colourless curved blades; melts at 266°. The β acid crystallises in stellate groups, melting at 250° to 252°. The α sodium salt crystallises with 4 mols. of water in colourless pointed blades; the β salt with 5 mols. of water in rhomboidal laminæ. The α barium salt crystallises with 7 mols. of water in long flexible needles; the β barium salt with 6 mols. of water in long brittle rectangular laminæ. In conclusion the author enters at some length into a theoretical consideration as to the constitutional formula of phenanthrene, which, in his opinion, consists of three benzene nuclei, one of which shares four adjacent carbon atoms with the two others—one ortho pair with each,—and phenanthrene may thus be derived from naphthalene by a repetition of the process by which the latter hydrocarbon is derived from benzene.

After some remarks by Dr. ARMSTRONG, to which Dr. JAPP briefly replied,

The SECRETARY read a paper "On some Derivatives of Phenyl-acetic Acid," by P. PHILIPPS BEDSON. This paper contains an account of the separation of para- and ortho-nitro-phenyl-acetic acids, both of which are crystalline substances; the former melts at 150° to 151°, the latter at 137° to 138°. Also of para- and ortho-bromo derivatives, which crystallise from water in white needles; the former melts at 114° to 115°, the latter at 103° to 104°. A di-bromo-phenyl-acetic acid has also been prepared by the long-continued action of bromine on mono-bromo-phenyl-acetic acid in sunlight; it crystallises in white needles, melting at 114° to 115°. A more detailed account is given of the bromo-nitro-phenyl-acetic acids and the corresponding brom-amido acids, short notices of which have appeared in the *Berichte Deuts. Chem. Gesell.*, x., 530 and 1065. A third isomeric acid has been obtained with these substances by nitrating crude bromo-phenyl-acetic acid. This β -bromo-nitro-phenyl-acetic acid crystallises from glacial acetic acid in small transparent prismatic needles, melting at 162°; yielding by reduction β -brom-amido-phenyl-acetic acid, melting at 186°.

The Society then adjourned to December 18, when the following paper will be read:—"On the Specific Volume and Density of Water of Crystallisation," by T. E. Thorpe. A ballot for the election of Fellows will also be held.

NOTICES OF BOOKS.

Blowpipe Analysis. By J. LANDAUER. Authorised English Edition, by J. TAYLOR and W. E. KAY. Pp. x. and 161. London: Macmillan, 1879.

We think that this is, on the whole, distinctly the clearest, most systematic, and most useful students' laboratory book on the employment of the blowpipe in chemical and mineralogical analysis. It includes the determination of artificial as well as natural inorganic compounds, and so differs from many other treatises on the blowpipe. The arrangement of the reactions obtained in tabular forms admitting of immediate inspection is carried to a considerable degree of completeness in this compact volume. A good many novelties in the way of apparatus and reagents are included amongst the descriptions, and take their place duly in the systematic course. Such additions have been inserted, however, only after repeated trials have proved their utility; so here are to be found Ross's aluminium plate support and Fletcher's miniature self-acting blowpipe; the employment of gold granules for detecting nickel and cobalt; and the use of hyposulphite of sodium and oxalic acid as a mixture for obtaining in the dry way characteristic metallic sulphides. Here are descriptions of flame-colourations as produced by Bunsen's methods; some condensed instructions as to spectrum analysis are also given. The book, indeed, so far as its object and scope allowed, is complete as well as systematic. Its contents naturally group themselves into five chapters, the first

treating of blowpipe apparatus and reagents, and the second of the actual operations and reactions in blowpipe testing. Then succeeds a chapter entitled "Special Examination for certain Elements in Combination," each element being studied in succession in its behaviour with the several tests. Chapter IV. gives a systematic course of analysis, while the remainder of the volume is devoted to the gathering up of the various experiments, observations, and inferences noted in preceding chapters into the form of tabular statements, and into condensed views of the behaviour of certain groups of compounds.

The book is carefully printed: indeed, we have detected but one misprint, ludicrous enough but of no importance, to be found on page 27, where the flame colouration of selenium is given as "corn flour blue" instead of "corn flower blue."

Report on a Preliminary Investigation of the Properties of the Copper-tin Alloys, made under the Direction of the Committee on Metallic Alloys. United States Board, to test Iron, Steel, and other Metals. By R. H. THURSTON, Chairman in the Mechanical Laboratory of the Stevens Institute of Technology. Washington: Government Printing Office.

WIDELY as metallic alloys have been employed in the arts, and frequently as they have been made the subjects of scientific research, our knowledge of them is still but fragmentary. Great merit, therefore, belongs to the United States Board for testing metals, and to their Committee on metallic alloys for undertaking the extensive and thorough going series of investigations, of which merely an instalment is recorded in the present volume.

The task undertaken by Prof. Thurston and his colleagues is very clearly defined. They have operated not on alloys formed of chemically pure materials and combined with all the precautions of the laboratory, but on such as are made of ordinary commercial metals by procedures practicable on the large scale. If we remember how small a proportion of some foreign bodies may seriously modify the nature of an alloy, we must admit that they have taken the right course for a series of experiments whose aim was purely practical.

With the chemical properties of the alloys, such as their powers of resistance to air, water, corrosive vapours, &c., they have not concerned themselves. The points determined have been the resistance of the alloys to transverse, tensile, torsional, and compressive stress. It has been ascertained that a comparatively limited group of compounds, extending from gun metal on the one hand to Muntz metal on the other, includes all that are really valuable. Alloys containing more than 27.5 per cent of tin are valueless for any purposes in which strength is requisite. The results of all the experiments are brought very clearly under the eye in a series of 21 graphic illustrations. Another important feature of the work is a set of photographs representing the ends of the fractured bars of metal, and showing more clearly than could be done by words the changes they have undergone from the strains applied. As an appendix there is a selection of papers by former authorities, both on the chemical and mechanical properties of alloys, and a list of the writers quoted.

Prof. Thurston and his coadjutors are engaged with similar researches on the alloys of copper and zinc, and on the triple compounds of copper, zinc, and tin. On the great utility of this report there can be no necessity for us to enlarge.

Crystalline Chlorophyll.—A. Trécul.—Referring to the paper of M. Arm. Gautier (*Bulletin de la Soc. Chimique*, xxviii., p. 147), the author states that as early as 1865 (*Comptes Rendus*, lxi., pp. 435, 436), he described such green crystals which he had seen formed directly from granules of chlorophyll.—*Comptes Rendus*,

CORRESPONDENCE.

ANALYSIS OF BOILER FEED-WATERS.

To the Editor of the Chemical News.

SIR,—In the CHEMICAL NEWS, vol. xxxix., p. 5, there appeared a scheme for the analysis of boiler feed-waters, by Mr. W. F. K. Stock, which commended itself to me as being a great advance upon the old method of giving the "temporary" and "permanent" hardness, and as being much more rapid and practical than the tedious method of resorting to a complete mineral analysis. There is, however, one important statement made, which I have not seen elsewhere, viz., that "*Calcium sulphate is practically insoluble at a pressure of one atmosphere of steam,*" and consequently should be debited to the salts precipitable in a steam boiler. Permit me to ask Mr. Stock, through the medium of your columns, if he will favour me with the nature of any evidence he may have at hand in support of this?

Naturally one would resort to the analysis of the scale from a boiler fed with water rich in SO_4Ca for information on this point, but unfortunately I have not such at hand. I may, however, mention that some time ago I found the incrustation from a tea-kettle to yield CO_2 equal to 94·16 per cent of CO_3Ca ; and some scale from a steam boiler gave $\text{CO}_2=71\cdot44$ per cent and $\text{SO}_4\text{Ca}=3\cdot66$ per cent only, the rest being chiefly organic matter and sand. The water producing the above contains about 8 grains of carbonate and 4 of sulphate of lime, but the boiler-water was taken from a river, hence the dirt it contained.—
I am, &c..

J. BACON.

Derby, December 4, 1879.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

*Comptes Rendus Hebdomadaires des Séances, l'Académie
des Sciences. No. 21, November 24, 1879.*

Heat of Formation of Ammonia.—M. Berthelot.—The author remarks that the heat of formation of ammonia, that of water, of carbonic acid, and of hydrochloric acid constitute the four most important data of thermochemistry. The three latter may be regarded as approximately known. For the formation of ammonia, dissolved and gaseous, he adopts the numbers $+21^{\circ}$ and $+72^{\circ}$.

Researches on Nitrification.—T. Schloesing and A. Muntz.—The authors have shown that natural nitrification may be considered as the result of a phenomenon analogous to fermentations, but that the oxidation of the nitrogen is not produced in a general manner by the organisms which are the common agents of the combustion of organic matter, but must be ascribed to a special organism. They have taken dilute alkaline solutions containing the necessary mineral matters, an ammoniacal salt, and organic matter. These liquids, after being heated under suitable conditions to 110° , remained without change for an unlimited time. But on allowing the access of atmospheric oxygen the formation of nitrates set in, and on examination with the microscope minute organisms were detected. The nitric ferment does not possess the power of resistance observed in some of its congeners. It is killed by exposure for ten minutes to a temperature of 100° . Sewage and similar liquids are generally rich in the nitric ferment, which assists, therefore, in their purification. It is also very abundant in vegetable earth, and generally in arable land.

Constitution of Dibromated Ethylen.—E. Demole. —The author seeks to establish the place of the two atoms of bromine with a view of deciding whether this compound is symmetric or asymmetric.

New Method for the Accurate Analysis of Commercial Potash.—B. Corenwinder and G. Contamine.—The authors add to the portion taken for analysis a slight excess of hydrochloric acid, and then, without taking any notice of the possible presence of sulphuric acid, silica, or phosphoric acid, they evaporate in the water-bath, after having added a sufficiency of platinum bichloride. The potassium chloro-platinate thus obtained is digested with alcohol at 95°, mixed with ether, and washed in the ordinary way with the same liquid. When this operation is complete, boiling water is poured upon the filter by means of a pipette, till the chloro-platinate is entirely dissolved, and the filtered liquid is collected. Water containing sodium formiate is then heated, and whilst it is boiling the preceding solution of potassic chloro-platinate is carefully poured into it by degrees. In a few moments the platinum is thrown down as a black powder, which merely needs to be washed, dried, heated to redness, and weighed in order to find the quantity of potassa present in the sample.

Bulletin de la Société Chimique de Paris,
Nos. 8 and 9, Nov. 5.

Various Thermo-Chemical Data.—M. Berthelot.—The author has determined the formation heat of cyanogen and of diamylen, the heat of fusion of glycerin and the specific heat of glycerin.

The Elimination of Bromine from Bromo-citraconic Acid and a New Organic Acid.—M. E. Bourgoin.—Already noticed.

Solubility of Benzoic and Salicylic Acids.—M. E. Bourgoin.—At 0° one part of benzoic acid requires for solution 588 parts of water, and 1 part of salicylic acid requires 666 parts.

The rest of this issue consists of extracts from other journals, which have been or will be noticed under their respective heads.

No. 10, December 5, 1879.

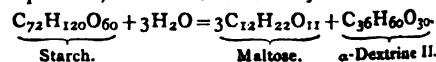
Hydrochlorate of Hydrogen Phosphide.—J. Ogier.
—The author has obtained this compound by compressing and refrigerating a mixture of equal volumes of the two gases.

Heat of Formation of Hydrobromate and Hydriodate of Hydrogen Phosphide.—J. Ogier.—Already noticed.

The Pyridic Bases.—A. Richard.—The bases which predominate in Dippel's oil are pyridin and lutidin; their joint proportion amounts to 40 per cent of the crude alkaloïds. Picolin and collidin are present in smaller quantities. A small quantity of ethylic alcohol has also been obtained from the crude oil.

Partial Synthesis of Milk-Sugar and Contribution to the Synthesis of Cane-Sugar.—E. Demole.—Already noticed.

Products of the Transformation of Starch.—C. O'Sullivan.—With reference to the paper of M.M. Macculis and Gruber (*Bulletin*, xxx., p. 54), the author remarks that they have doubtless overlooked his memoir in the *Journal of the Chemical Society* (ii., p. 125, 1876), in which he has shown that the molecular scission of starch takes place not according to one single equation, but to three equations, to which a fourth may be added:—



On Chlorophyll.—A. Gautier.—Already noticed.

Dingler's Polytechnisches Journal.
Band 234, Heft 4.

Production and Application of Phosphorescent Powders.—The patentees of this process (Prince Sagan, W. F. McCarty, and E. Peiffer) employ a mixture of 100 parts carbonate and phosphate of lime (obtained by the ignition of shells, especially *Tridama* and *Sepia*) with 100 parts quicklime, 25 parts of calcined salt, and 25 to 50 per cent of the whole mass of sulphur: 6 to 7 per cent of a colouring-matter—a sulphide of calcium, strontium, barium, magnesium, aluminium, &c.—must then be added. This powder serves to render barometers, compasses, &c., luminous, and is particularly phosphorescent under the influence of an electric current.

On the Leblanc Soda Process.—F. Fischer.—The author has examined the composition of the gases escaping from the black-ash furnaces, and the temperature of the melting mass. On one occasion only were dubious traces of carbonic oxide detected.

Acidimetry of Coloured Acids.—Dr. F. Salomon.—The author has devised an apparatus for the determination of crude wood-vinegar and other highly-coloured products. The process cannot be intelligibly described without the accompanying illustration.

Products of the Fats obtained by Acid Saponification on Distillation with Superheated Steam.—The fatty acids produced include the valerianic, capronic, ceanthylic, and caprylic.

Preparation of Red and Brown Naphthalin Dyes.—An abstract of the German Patent No. 5411, March 12, 1878.

Development of Dyeing, Printing, and Bleaching. Dr. A. Kielmeyer.—The continuation of a lengthy paper, not capable of useful abstraction.

On Methyl-anilin and Methylic Alcohol.—M. Bardy.—A summary of the methods for the production of these compounds, and for the detection of ethylic alcohol in admixture with the methylic.

Chemiker Zeitung.
No. 45, 1879.

Considerable sensation has been excited by the report that arsenic had been detected in the paper collars, &c., manufactured by a Leipzig firm. On a careful examina-

tion, conducted by six of the most eminent chemists, the accusation was proved to be utterly unfounded.

The explosion of a calendering machine at Gera, on November 21st, proved fatal to four persons.

MISCELLANEOUS.

The Science and Art Department.—From the Twenty-sixth Report of the Science and Art Department of the Committee of Council on Education, we learn that during 1878 there were 36 regular and 189 occasional students at the Royal School of Mines. At the Royal College of Chemistry there were 303 students, and at the Metallurgical Laboratory 74. The lectures delivered in the Lecture Theatre at South Kensington were attended by 5491 persons. The evening lectures to working men at the Royal School of Mines were attended by 1685 persons, being 458 more than in 1877; and 169 science teachers attended the special courses of lectures at the Science Schools, South Kensington. At the Royal College of Science for Ireland there were 22 associates or regular students, and 53 occasional students; while the various courses of lectures there were attended by 1421 persons.

MEETINGS FOR THE WEEK.

SATURDAY, 13th.—Physical, 3. "A New Form of Resistance Balance for Comparing Standard Coils," J. A. Fleming, D.Sc. "The Graduation of Prof. Hughes' Sonometer," J. H. Poynting. "A Dispersion Photometer," W. E. Ayrton and J. Perry. "The Value of 'g' at Tokio, Japan," W. E. Ayrton and J. Perry.

MONDAY, 15th.—Medical, 8.30.
London Institution, 5.
Society of Arts, 8. (Cantor Lecture). "Chemistry of Bread and Bread Making," Dr. Graham.

TUESDAY, 16th.—Civil Engineers, 8.
Zoological, 8.30.

WEDNESDAY, 17th.—Society of Arts, 8. (Ordinary Meeting). "The Panama Canal," Captain Bedford Pim, R.N.
Geological, 8.
Meteorological, 7.

THURSDAY, 18th.—Royal, 8.30.
Chemical, 8. Ballot for the Election of Fellows. "On the Specific Volume and Density of Water of Crystallisation," T. E. Thorpe. "On the Analysis of Organic Bodies Containing Nitrogen &c.," W. H. Perkin.
Philosophical Club, 6.30.

COMPOSITION AND QUALITY OF THE METROPOLITAN WATER.

NOVEMBER, 1879.

The following are the returns of the Society of Medical Officers of Health:—

	Appearance in a foot Tube.	Ammonia.		Nitrogen as Ni- trates, &c.	Oxygen used to Oxidise Organic Matter.	Total Solids.	Lime.	Magnesia.	Chlorine.	Carbonic An- hydride.	Hardness on Clark's Scale		
		Saline. Grains.	Organic. Grains.								Before Boiling. Degrees	After Boiling.	
<i>Thames Water Companies.</i>													
Grand Junction	Clear	0.000	0.009	0.180	0.050	21.10	7.630	0.576	1.152	1.400	14.3	3.7	
West Middlesex	Clear	0.000	0.008	0.150	0.043	21.50	7.840	0.504	1.152	1.170	14.3	4.2	
Southwark and Vauxhall	Clear	0.000	0.008	0.132	0.048	22.10	7.340	0.360	1.152	1.330	14.8	3.7	
Chelsea	Slightly turbid	0.000	0.008	0.144	0.055	21.00	7.590	0.396	1.080	1.500	14.3	3.7	
Lambeth	Clear	0.000	0.009	0.144	0.062	21.50	7.050	0.504	1.152	1.270	13.2	3.3	
<i>Other Companies.</i>													
Kent	Clear	0.000	0.002	0.450	0.039	28.60	9.070	0.896	1.872	1.400	18.8	7.5	
New River	Clear	0.000	0.007	0.138	0.014	20.60	7.630	0.432	1.152	1.730	14.8	3.3	
East London	Clear	0.000	0.007	0.135	0.031	21.50	7.840	0.504	1.224	1.330	15.4	4.2	

The quantities of the several constituents are stated in grains per imperial gallon.

NOTE.—The amount of oxygen required to oxidise the organic matter, nitrates, &c., is determined by a standard solution of permanganate of potash acting for three hours.

C. MEYMOTT TIDY, M.B.

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LECTURE ARRANGEMENTS BEFORE EASTER, 1880.

LECTURE HOURS, THREE O'CLOCK. CHRISTMAS LECTURES.

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Professor EDWARD A. SCHÄFER, F.R.S.—Ten Lectures on The Physiology of Muscle; on Tuesdays, Jan. 13 to March 16. One Guinea.

Mr. HEATHCOTE STATHAM, Esq.—Two Lectures on Modern Architecture since the Renaissance; on Thursdays, Jan. 15 and 22. Half-a-Guinea.

Professor DEWAR, M.A., F.R.S.—Eight Lectures on Recent Chemical Progress; on Thursdays, Jan. 29 to March 18. One Guinea. Professor of 1. RUPERT JONES, F.R.S.—Three Lectures on Coal; on Saturdays, Jan. 17, 24, 31. Half-a-Guinea.

Professor of 2. ERNST PAUER.—Three Lectures on Handel, Sebastian Bach, and Joseph Haydn. With Musical Illustrations. On Saturdays, Feb. 7, 14, 21. Half-a-Guinea.

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The Friday Evening Meetings will begin on January 16th, at 8 p.m. Professor Dewar, F.R.S., will give a Discourse (Studies on the Electric Arc) at 8 p.m. Succeeding Discourses will probably be given by Dr. W. B. Carpenter, Professor J. Marshall, Dr. Huggins, Mr. W. H. Frazer, Rev. H. R. Haweis, Mr. F. J. Bramwell, Mr. H. N. Moseley, Dr. C. William Siemens, Professor Tyndall and Huxley, Lord Rayleigh, Mr. G. J. Romanes, M. Lecoq de Boisbaudran, Mr. H. Pollock, Professor Frankland, Mr. H. H. Statham, Mr. W. Spottiswoode, and Mr. Warren De la Rue. To these Meetings Members and their Friends only are admitted.

Persons desirous of becoming Members are requested to apply to the Secretary. When proposed, they are admitted to all the Lectures, to the Friday Evening Meetings, and to the Library and Reading Rooms; and their Families are admitted to the Lectures at a reduced charge. Payment—First Year, Ten Guineas; afterwards, Five Guineas a Year; or a composition of Sixty Guineas.

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ALEX. MOORE, Secretary.

136, Hope Street, Glasgow,
December 10th, 1879.

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THE CHEMICAL NEWS

VOL. XL. No. 1047.

THE PERIODIC LAW OF THE CHEMICAL ELEMENTS

By D. MENDELÉEF.

(Continued from page 260)

IN spite of the similarity of the corresponding compounds of Na, Cu, and Ag, the metals can be very easily distinguished one from another, which agrees with the fact that the volume of the metal alone is not the same as the volume of the metal when combined with another element. The volume of Na=24, of Cu=7, of Ag=10. The atoms of Na are further apart from each other and are more susceptible to reaction than the atoms of Cu and Ag. When Na enters into combination with another body, there is often a considerable contraction takes place. For example, the volume of Na=24, the volume of NaHo=19; 24 volumes of Na give 27 volumes of NaCl, whilst 10 volumes of Ag give 26 volumes of AgCl. This and many other examples show how arbitrary it is to conclude from the known volume of a compound the unknown volumes of its elementary parts.

We see, from the inspection of Table I.,* that the members at the beginning of the large periods (as well as the small periods commencing with Na and Li) are metals of a very strongly pronounced alkaline nature, whilst the members at the end are energetic haloids. Already, in the time of electro-chemists, the above-mentioned elements occupied the same positions of extreme members in the system. It is incontestable that the electro-chemical theory has had a great deal of influence on the later progress of chemistry. However, I am far from wishing to set myself up as a defender of this theory. To show this I will mention that, as can be seen in Table I., the first and last members of the large periods are the only ones which have a clearly defined chemical character. In other words, they are the only elements which react easily, and at temperatures which are not too high, with the greater number of other bodies; they alone possess atomic weights of about the same value, and some other analogous properties. The volume of a compound body increases nearly in equal quantities when H is replaced by Cl (=35.5), or by K (=39). When metaleptic hydrogen is replaced by chlorine the volume increases in a nearly constant ratio. For examples:—

Volume C_3H_{12} = 110	C_6H_6 = 87	C_7H_8 = 104
" $C_3H_{11}Cl$ = 117	C_6H_5Cl = 97	C_7H_7Cl = 110
Diff. of vol. when H —	—	—
is replaced by Cl 7	10	6

The volume increases in an analogous manner, and in nearly the same quantity, when H is replaced in the acids by K.

Volume HCl = 29	H_2O = 20	HNO_3 = 41	H_2SO_4 = 53
" KCl = 37	KHO = 28	KNO_3 = 48	K_2SO_4 = 66
Diff. of vol. when H is replaced by K	8	7	13

We can bring together, according to their atomic weights, the following unlike elements:—

O = 16	F = 19	Na = 23	Mg = 24
S = 32	Cl = 35.5	K = 39	Ca = 40
Se = 78	Br = 80	Rb = 85	Sr = 87
Te = 125?	I = 127	Cs = 133	Ba = 137

* See CHEMICAL NEWS, vol. xl, p. 268.

The transition from Cl to K, &c., corresponds also, in many respects, to a certain analogy between these elements, although in the same period there are no elements having such similar atomic weights and such different properties. Because of this last fact, it might be better to separate the series here, in such a manner that the large periods commence with K and terminate with Cl. But in reality the series of elements is uninterrupted, and represents in a certain degree a spiral function.

In consequence of this, the most dissimilar elements are found at the beginning and at the end of large periods; in the middle are the transitory elements, and the most analogous are the nearest together. It is in this manner that after the elements of the alkalies and the alkaline earths come these rare elements (metals of gadolinite and cerite, Ti, V, Cr, Nb, Mo, Ta, W, Ur), which, even from an analytical point of view, are similar amongst themselves. They are not volatile; they are difficultly fusible, and difficult to reduce; they possess, even in their highest oxides, a feeble power of reaction; they are often found together in nature, and rarely in large quantities, &c. These elements are rare, and I even think that that is because of their character and not by chance. We can mention as a comparison that among the hydrocarbides the series C_nH_{2n+2} and C_nH_{2n-6} are often found together in nature, both being formed in many reactions; they possess a pronounced reaction and furnish many derivatives, while the members of the intermediate series, particularly C_nH_{2n} and C_nH_{2n-2} are formed more rarely, and do not give so many independent combinations as the preceding series. Our knowledge about the rare elements that I have just mentioned is unfortunately very defective, and if we had not before us the classic researches of Marignac on the compounds of Zr, Nb, and Ta, the entire group would be a collection of elements without any importance in the system. The works of Blomstrand, of Roscoe, of Delafontaine, and of Bunsen have thrown much light on the characters of the rare elements; but these elements are still the object of a series of questions which have not yet been answered.

After these rare elements (see Table I.) comes first in importance the group of elements called noble; in the system they all grouped together. The useful metals are attached to them, and by the help of As, Sb, and Te, they lead on to the metalloids.

In the groups of analogous elements, the elements whose atomic weights are the highest possess more strongly marked basic properties, or form weaker acids. Bunsen has shown that Ca is more electro-positive than K and Rb; the basic properties are more apparent in BaO than in CaO, in ThO_2 than in ZrO_2 or TiO_2 ; HgO separates MgO or BeO from compounds; Bi_2O_3 is a more energetic base than Sb_2O_3 or As_2O_3 ; P_2O_3 does not exhibit any basic properties, unless it is in this circumstance, that PH_3O_3 is not a tribasic acid, but a dibasic acid. For the same reason Ta gives a more feeble acid than Nb and V, Te an acid more feeble than Se and S. The acid properties are very feebly marked in PbO_2 ,* although Frémy obtained a salt, $K_2PbO_3 \cdot 3H_2O$, corresponding to salts prepared by means of stannic and silicic acids; SnO_2 possesses feeble basic and acid properties; and SiO_2 cannot act except as an acid, although as a weak acid.

Besides this, when the atomic weight increases in the members of a determined group, it is not only that the faculty of being reduced to the state of a simple body increases (compare Te and Se with S, I with Cl, Au with Cu, &c.), but also the power of giving lower forms of oxidation, exhibits itself frequently by a relative stability and by a great aptitude for forming compounds. Bi forms Bi_2O_5 with difficulty—generally the compounds of bismuth correspond to Bi_2O_3 , or BiX_3 ; in the same manner

* A true analogue of SnO_2 , PbO_2 , should also be able to act as a base. However, the attempts that I have made, above all in that which concerns the action of HF, have led to no results. To all appearance the known form, but answers to metastannic and metatitanic acid. We must wait for the discovery of a variety of PbO_2 with basic properties.

Pb does not only form PbO_2 , but also the very stable oxide PbO , which Sn and Si are not capable of doing to the same degree; Tl does not only form Tl_2O_3 , but also Tl_2O , which has not been observed either in the case of In or of Al.

In the group Mg, Zn, Cd, Hg, we notice, with the augmentation of the atomic weight, a marked increase in the volatility, basic properties of the oxide RO, reducibility to the metallic state, and the power of forming a lower oxide, R_2O .^{*} The volatility does not increase with the augmentation of the atomic weight, except in this and the neighbouring series; on the contrary, it diminishes in the last series, as is well shown by Cl, Br, and I.

It is precisely for the motives expressed above that the so-called noble metals occupy the place which has been given to them in the system,—that is, in the middle of the large periods, amongst the members having high atomic weights, which comprises elements which are very easily reducible, and whose reactions are very weak.

All that has preceded shows the nature of the periodic law. No natural law acquires any scientific importance unless it introduces, so to speak, some practical conclusions, or, in other words, unless it admits of logical conclusions capable of explaining what has before remained unexplained, and, above all, unless it raises questions which can be confirmed by experience. In a case of this nature the use of the law is evident, and it is possible to control its correctness.

This law will at least incite research into the new, little-known parts of science. This is why I propose to study attentively some consequences of the periodic law, and to examine how it can be applied—

- To the system of elements;
- To the determination of the atomic weights of insufficiently studied elements;
- To the determination of the properties of elements up to the present time unknown;
- To the correction of atomic weights;
- And to the enlargement of our knowledge of the forms of chemical combinations.

I shall not form any hypotheses, either here or further on, to explain the nature of the periodic law; for, first of all, the law itself is too simple;† and, secondly, this new subject has been too little studied yet, in its diverse parts, for us to be able to form any hypothesis: the most important reason is the third one,—that is, that we cannot put the periodic law and the atomic theory in accord without upsetting the known facts about the exact values of the most carefully found atomic weights. There is, however, I believe, between the series of elements and the homologous series, an analogy which, although small, is not less real, which authorises me to admit that it is so; it is the comparison of the physical properties of a group to which I shall return later on.

(To be continued.)

THE DISSOCIATION OF CHLORINE.

VICTOR MEYER'S ANSWER TO SEELHEIM'S CRITIQUE.‡

SINCE I communicated to you Seelheim's experimental critique§ of V. Meyer's famous experiments this investigator has published an answer to that critique, which (I rejoice to say) amounts to a complete and (I believe) unanswerable refutation.

* It can be foreseen that cadmic oxide will give a basic suboxide, Cd_2O ,—certainly very unstable in air,—or else salts CdX , which correspond to it.

† However, I do not ignore that to completely understand a subject we should possess, independently of observations (and experiences) and of laws (as well as systems), the meanings of both one and the other.

‡ From an article in the *Ber. der Deutsch. Chem. Ges.*, No. 18. Communicated by Prof. Dittmar.

§ *CHEMICAL NEWS*, vol. xl., p. 244.

What V. Meyer says is easily condensed in a few words. It is quite true that at a "yellow" heat platinum-metal can be volatilised in a current of chlorine. V. Meyer himself has often repeated Troost and Hautefeuille's experiments; but, in looking into the matter quantitatively, he found that at $-1370^{\circ}C$. a given quantum of metallic platinum, when exposed to a rapid current of dry chlorine for a whole hour, lost only 1 per cent of its weight. In his vapour-density determinations the platinum produced (from the $PtCl_4$) is exposed to a stagnant atmosphere of chlorine for only a few seconds. It is difficult to assume that, under these circumstances, more than perhaps a few hundredths of a per cent of the metal could possibly get gasified. But, what is more, he always took care, after the experiment, to examine the wee bucket which served for the introduction of the platinous chloride, and he invariably found it full of spongy metal, the weight of which (whenever the test was applied) agreed almost absolutely with that calculated from the weight of the chloride used. Of crystalline or sublimed platinum no trace could be discovered.

Moreover, iodine, as V. Meyer stated in his first communication, behaves at high temperatures as chlorine does, although it naturally was used as such, and not in the shape of a platinum compound at all.

It appears, then, that V. Meyer after all is right, and that we must admit that the substance Cl_2 at high temperatures suffers decomposition, perhaps according to equation $2Cl_2 = Cl_2 + Cl + Cl$. This from the first was my theory of the phenomenon, and I mean to retain it until it is disproved. Not that I claim any credit for an idea which is so obvious that it must have suggested itself to any chemist who took the trouble of reasoning on the matter.

Lieben* (besides offering this very obvious explanation) suggests that possibly V. Meyer's result may be owing only to the fact that the coefficient of expansion of chlorine increases rapidly at high temperatures, and he does not see that he thereby only re-states the problem. In any gas of moderate tension the molecules must be assumed to be mere points compared with their average distances. If such a thing expands on being heated it can do so only through two causes, viz., first, an increase in the energy of the progressive motion of the molecules; and secondly, a multiplication of molecules, i.e., through dissociation. The expansion owing to the first cause amounts to 1/273rd of the volume at $0^{\circ}C$ per degree Centigrade. To say that a certain gas, within a certain interval of temperature, expands at a greater rate is only another way of stating that the number of molecules in it increases in consequence of the respective elevation of temperature.

W. D.

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SEPARATION OF PHOSPHORIC ACID FROM THE SESQUIOXIDES OF IRON AND ALUMINA.

By M. P. DEROME.

THE separation of phosphoric acid when combined with sesquioxides of iron and alumina is effected in a very satisfactory manner by the following method:—

The substance, mixed with from five to six times its weight of dry sodic sulphate, is strongly heated for eight or ten minutes over an enameler's blast-lamp; when cool the mass is extracted with water, which dissolves the excess of sodic sulphate and the phosphoric acid in the state of tribasic phosphate of soda.

The phosphoric acid in this liquid can be determined either by means of a standard solution of uranium, or by

* Extract from *Comptes Rendus* No. 6 in No. 18 of the *Berichte*.

precipitation of the phosphoric acid in the state of argentic phosphate or ammonio-magnesian phosphate.

This method may be applied to the treatment of phosphoric acid in soils, iron ores, and generally in every material containing small quantities of phosphoric acid and much iron and alumina.

In soils phosphoric acid compared with the sesquioxides is always found in feeble quantities. It suffices, then, after the elimination of the silica and of the residue insoluble in acids, to search for phosphoric acid in the precipitate of ferric oxide and aluminium, as obtained by the aid of heat from a solution almost neutral in presence of sodic acetate.

If a very calcareous soil does not contain sufficient sesquioxides for the total precipitation of phosphoric acid, a slight excess of ferric oxide may be added; the richness of the precipitate in phosphoric acid is a sure indication of the sesquioxides being in sufficient quantity.

Synthetical experiments have given amongst others the following results, which sufficiently show the value of the method:—

		Phosphoric Acid	
Fe ₂ O ₃	Al ₂ O ₃	Introduced.	Found.
Introduced.	Alumina Oxide.		
0.308	0.108	0.025	0.0249
0.195	0.300	0.010	0.010
0.195	0.300	0.040	0.040
0.310	0.115	0.125	0.1247
0.110	0.300	0.125	0.125
0.310	0.115	0.005	0.005
0.110	0.300	0.005	0.005

—Comptes Rendus, December 1st.

FURTHER EXPERIMENTS ON THE VAPOUR DENSITIES OF POTASSIUM AND SODIUM.*

By Professor JAMES DEWAR, M.A., F.R.S., and
ALEXANDER SCOTT, B.A.

In our former communication "On the Vapour Densities of Potassium and Sodium,"† we pointed out the chief obstacles which lay in the way of an exact determination of these constants. Having overcome the chief manipulative difficulties in connection with the method we described, there still remained the problem for solution as to how far the use of iron bottles in our experiments might affect the results. If the iron retained the metals or allowed their vapours to diffuse with rapidity through it, a considerable error might be produced without its being easily detected. We noted an action of this kind, remarking that "both metals seemed to have a tendency to behave in the same manner as zinc, viz., to form an alloy with the iron, because the vapour evolved after a few minutes was absorbed completely, in spite of all precautions to prevent oxidation." The tendency of iron and the alkali metals to form alloys, we thought, would be diminished, so as not to interfere with the value of the results, "by raising the temperature as much as possible, so as to induce rapid volatilisation," and the fair accordance of the series of experiments formerly given led us to suppose that this difficulty had been successfully obviated. Recent spectroscopic observations, by Professors Leving and Dewar, on the luminous absorption of the vapours of sodium under different conditions in iron and platinum vessels—which will be presented to the Society—having led to the inference that mere rapidity of volatilisation, of such small weights of the metals as can be conveniently employed for the measurement of the volumes of the vapours, in presence of a relatively enormous mass of iron, would not eliminate the absorptive action, new experiments were made in a platinum apparatus of similar shape and size to the iron one, and in the same manner as formerly described. Magnesia was used instead of sand to pack

the platinum vessel in the crucibles, employed as a protection against the diffusion of furnace gases, and the volatilisation of the alkali metals took place in nitrogen. Each set of experiments was made at different times, and the results are given in the order of observation, the bottle being freed from the vapours after each experiment by a strong current of hydrogen. The molecular weights are calculated for 22.34 c.c. at 0° C. and 760 m.m. pressure.

The following tables contain the record of the results obtained:—

Potassium.

I.

0.046 grm. gave	24.1 c.c. at 0° and 760 m.m.	Mol. wt.	42.6
0.045 "	23.1 "	"	43.5
0.042 "	21.2 "	"	44.2
0.045 "	22.9 "	"	43.9
0.045 "	22.1 "	"	45.5
0.030 "	25.1 "	"	44.5

Mean molecular weight 44.03

II.

0.047 grm. gave	26.8 c.c. at 0° and 760 m.m.	Mol. wt.	39.2
0.049 "	26.0 "	"	42.1

Mean molecular weight 40.65

Sodium.

I.

0.032 grm. gave	26.5 c.c. at 0° and 760 m.m.	Mol. wt.	27.0
0.033 "	26.7 "	"	27.6
0.031 "	27.9 "	"	24.8

Mean molecular weight 26.4

II.

0.032 grm. gave	25.1 c.c. at 0° and 760 m.m.	Mol. wt.	28.5
0.032 "	26.3 "	"	27.2
0.031 "	27.9 "	"	24.8
0.029 "	26.5 "	"	24.4
0.031 "	26.9 "	"	25.7
0.030 "	27.3 "	"	24.6
0.030 "	26.1 "	"	25.6

Mean molecular weight 25.8

III.

0.027 grm. gave	24.6 c.c. at 0° and 760 m.m.	Mol. wt.	24.5
0.027 "	23.2 "	"	26.0
0.037 "	31.7 "	"	26.1
0.030 "	29.1 "	"	23.0

Mean molecular weight 24.9

The new determinations of the molecular weights of potassium and sodium are just about half the former values, and would seem to support the inference that the atom of each of these metals resembles mercury and cadmium in the gaseous state, as regards molecular volume. Such a remarkable result cannot be accepted without a very thorough investigation of the secondary reactions when any take place, under the conditions of the experiment. It is certain that in the platinum vessels there is much less absorption of the vapour than in the case of iron, and that the action of the furnace gases is all but eliminated. The following suggestion towards an explanation of the results naturally occur. The metals may contain a large amount of occluded hydrogen. With reference to this assumption, we have examined the sodium used in our experiments and find, on careful exhaustion and boiling in a Sprengel vacuum, that one volume of sodium contains only one-third of its volume of occluded hydrogen, so that this will not account for the double volume of the atom of

* Abstract of Paper read before the Royal Society
† Proc. Roy. Soc., January, 1879.

the metals." The alkali metals may form volatile compounds with platinum, although, judging from analogy, such compounds are not likely to exist. If so, the vapour blown out of the bottle ought to contain platinum. The use of platinum tubes to blow out the vapours prevented the test being applied. Occluded oxygen from the platinum might transform the metals into oxides, and thus complicate the results. We have, however, always observed plenty of free metallic vapour on blowing out the bottle, so that the metals remain chiefly in the state of vapour. Unless some other explanation can be given, it will be necessary to admit some anomaly in the behaviour of potassium and sodium, as regards their vapour densities at high temperatures. In future the variation of the density with the temperature must be determined, but this is very difficult at temperatures about 1500° C. The chief drawback to the prosecution of the experiments is the terrible waste of platinum vessels, which never stand the combined action of potassium, sodium, and hydrogen at high temperatures beyond two or three operations.

The following determinations of the molecular weight of potassium iodide, made in the iron apparatus, are of importance in connection with the previous results.

Potassium Iodide.

0.159 grm.	gave	20.75	at	0°	and	760 m.m.	Mol. wt.	171.0
0.165	"	21.2	"	"	"	"	"	174.0
0.170	"	21.4	"	"	"	"	"	177.0
0.180	"	21.6	"	"	"	"	"	186.0
0.193	"	15.1	"	"	"	"	"	152.0
0.174	"	24.4	"	"	"	"	"	159.0

Mean molecular weight 169.8

These numbers are fairly accordant, and seem to indicate that iodide of potassium is normal in its behaviour. The vapour after each experiment was blown out with a current of hydrogen. A crystalline deposit was obtained in each case, which was pure iodide of potassium, free from any trace of iron. Taken in connection with the former experiments, this seems to show that, if free potassium is abnormal, its compounds are not altered. Before any final conclusions can be reached, further experiments must be made in platinum or iridium vessels, and it will be very important to control the results by examining the densities of the iodides of caesium and rubidium.

ON THE VAPOUR-DENSITIES OF
PEROXIDE OF NITROGEN, FORMIC ACID,
ACETIC ACID, AND PERCHLORIDE OF
PHOSPHORUS.

By J. WILLARD GIBBS.

(Concluded from p. 285.)

In Table VI. are exhibited the results of experiments by Playfair and Wanklyn,* in which the vapour of the acid was diluted with hydrogen or, in a single case (the experiment at 95.5°), by air. Columns I. and II. of the observed densities relate each to a series of observations by the method of Gay-Lussac; column III. contains four independent determinations by the method of Dumas. The numbers in the column of pressures are, as in other similar cases, the partial pressures obtained by subtracting from the total pressure (which was never very much less than that of the atmosphere) that which would be exerted by the hydrogen or air alone.

The first observation of the first series gives the density 1.036, which is doubtless too small, since it is much less than the theoretical limit 1.073. Since the greater part of the measurements from which this number was calcu-

lated were also used in reducing the other observations in the series the error probably affects the other observations and in a somewhat increased degree. This will account only for a part of the difference between the observations and the formula. The remaining part of the differences in this series, and the somewhat smaller differences in the next, may be due to the fact that the experiments of both series were conducted with descending temperatures. Yet the experiments of the third column, which were made by Dumas's method, do not exhibit any preponderance of positive values for the excess of observed density, but rather the opposite.

On the whole, these experiments furnish no decisive indication of any influence of the hydrogen or air upon the vapour. They may be thought to corroborate slightly the tendency observed in the experiments of Naumann and Troost to lower densities than the formula gives at very low pressures. Yet where the experiments of Naumann show the greatest deficiency in observed density (at 78° and 80 m.m.), an experiment of Playfair and Wanklyn, at almost the same temperature and pressure, gives a trifling excess of observed density, and at a little lower temperature and pressure, where we should expect from the experiments of Naumann that the deficiency would be still greater, an experiment of Playfair and Wanklyn shows a great excess of density.

By combining the experiments of Cahours, Naumann and Troost, we may obtain observations of density at 130° for a very wide range of pressures. For one atmosphere, we may regard the formula as coinciding with the average of the numbers given by Cahours. For pressures between three-quarters and one-half of an atmosphere the experiments of Naumann show an excess of density; at pressures below half an atmosphere the experiments both of Naumann and of Troost show a deficiency of density as compared with the formula. For an indefinite diminution of pressure there can be little doubt that the real density, like the value given by the formula, approaches the theoretical value 1.073. The greatest excess in the numbers obtained by experiment is 0.07; the greatest deficiency is 0.19, which occurs at 59.7 m.m. The next in order of magnitude is 0.11, which occurs more than once. These discrepancies are certainly such as may be accounted for by errors of observation. They do not appear to be greater than we might expect on the hypothesis of the entire correctness of the formula. On the other hand, the agreement is greater than we should expect, if we reject the theory on which the formula was obtained. It is about such as we might expect in a suitable formula of interpolation with three constants, which have been determined by the values of the density for one atmosphere, for half an atmosphere, and for infinitesimal pressures. But we must regard the actual formula, in its application to this single temperature, as having only two constants, of which one is determined so as to make the formula give the theoretical value for infinitesimal pressures, and the other so as to make it agree with the experiments of Cahours at the pressure of one atmosphere.

An entirely different method has been employed by Horstmann* to determine the vapour-density of this substance. A current of dry air is forced through the liquid acid, which is heated to promote evaporation, and the mixture of air and vapour is cooled to any desired temperature, with deposition of the excess of acid, by passing upward through a spiral tube in a suitable bath. The acid is then separated from the air, and the quantity of each determined. It is assumed that the air is exactly saturated with vapour on leaving the coil, and that it has the temperature of the bath. If we know the pressure of saturated vapour for that temperature, and assume the validity of Dalton's law, it is easy to calculate the density of the vapour. For the pressure of the air is found by subtracting the pressure of the vapour from the total pressure (the experiments were so conducted that this was the same as

* *Trans. Roy. Soc. Edin.*, vol. xiv. p. 466.

* *Berichte der Deutschen Chemischen Gesellschaft*, Jahrg. iii. 1870, s. 78; and Jahrg. vi. (1873) s. 280.

TABLE VI.—ACETIC ACID.
Experiments of Playfair and Wanklyn.

Tempera- ture.	Pres- sure.	Density calc. by eq. (12).	Density observed.			Excess of observed Density.		
			I.	II.	III.	I.	II.	III.
212.5	322.8	2.124		2.060			-.064	
194	326.0	2.168		2.055			-.113	
186	334.4	2.173	1.936			-.237		
182	319.4	2.213		2.108			-.105	
166.5	289.5	2.293		2.350			+.057	
163	245.8	2.290	2.017			-.273		
132	227.5	2.628	2.292			-.336		
130.5	205.7	2.729		2.426			-.303	
119	269.0	2.914		2.623			-.291	
116.5	211.3	2.876	2.371			-.505		
95.5	(123.8)	3.105		2.594			-.511	
86.5	(200.4)	3.432		3.172			-.260	
79.9	(83.3)	3.297		3.340			+.043	
62.5	(46.2)	3.473		3.950			+.477	

the actual pressure of the atmosphere), and the ratio of the weights of the acid and the air obtained by analysis, divided by the ratio of their pressures, will give the ratio of their densities. The pressures of saturated vapour employed by Horstmann are those given by Landolt,* and differ greatly from the determinations of Regnault, in some cases being nearly twice as great,—a difference noticed but not explained by Landolt, who, however, gives determinations (previously unpublished) of Wüllner, which somewhat exceed his own. (On the other hand, the observations of Bineau substantially agree with those of Regnault.)

If we compare the observations of Horstmann with the values given by equation (12) on the basis of Landolt's pressures we find a very marked disagreement, as may be seen by the following numbers, which relate to the highest temperatures of Horstmann's experiments, where the disagreement is least:—

Temperature—								
63.1	62.9	59.9	51.1	49.0	48.7	44.6	41.4	
Pressure (Landolt)—								
110.0	109.2	97.0	69.0	63.4	63.0	53.1	46.6	
Density calculated by eq. (12)—								
3.67	3.67	3.69	3.75	3.77	3.77	3.79	3.81	
Density observed—								
3.19	3.11	3.12	3.16	2.89	2.98	2.75	2.62	

It will be observed that while the values obtained from equation (12) increase with diminishing temperatures the values obtained from Horstmann's experiments diminish. This diminution continues as far as the experiments go, until finally at 12° or 15° the densities are only one-half as great as those obtained by Bineau, by direct experiment at the same temperatures and at somewhat less pressures, in a series of observations which bear every mark of a very exceptional precision. (Compare Tables VII. and IV.) The explanation of this disagreement is doubtless to be found in the values of the pressures employed in the calculations, and it will be interesting to see how the results may be modified by the adoption of different pressures.

In determinations of the pressure of saturated vapours too great values are so much more easily accounted for than errors in the opposite direction, especially when the pressures are small, that especial interest attaches to the lowest figures which are supported by a competent authority. The experiments of Regnault were made with three different preparations of acetic acid, of which the second was once, and the third twice, purified by distillation over anhydrous phosphoric acid. Each distillation considerably diminished the pressure of the saturated vapour, the effect of the second distillation being about

half that of the first. The numbers obtained with the third preparation are given in the following table, with their logarithms and the differences of the logarithms for one degree of temperature:—

Temperature.	Pressure.	Log. Pressure.	Diff. per 1°.
9.71	6.42	0.8075	0.0239
12.12	7.33	0.8651	0.0272
14.33	8.42	0.9253	0.0161
14.87	8.59	0.9340	0.0252
17.23	9.85	0.9934	0.0251
19.84	11.455	1.0590	0.0237
22.37	13.15	1.1189	0.0232
25.28	15.36	1.1864	

The uniformity of the numbers in the last column shows the remarkable precision of the determinations. At the same time it is evident that the differences in these numbers are due principally to the errors of observation, so that numbers obtained by interpolation between the logarithms of the observed pressures will be somewhat better (on account of averaging of the errors) than the original determinations.

The values obtained by such an interpolation have been used for the comparison of Horstmann's experiments with the formula (12) which is given in Table VII. Unfortunately, this comparison cannot be extended above 25°, which is the limit of Regnault's experiments. The first three columns of the table give the temperatures of Horstmann's experiments, the pressures corresponding to these temperatures according to the determinations of Landolt, and the density deduced from Horstmann's experiments by the use of these pressures. To these columns, which are taken from Horstmann's paper, are added the pressure derived from Regnault's observations by the logarithmic interpolation described above, the density calculated by equation (12) from these pressures and the temperatures of the first column, and the densities obtained by combining Horstmann's experiments with Regnault's pressures. This column is derived from the second, third, and fourth, as follows. If w and W denote respectively the weights of vapour and of air which pass through the apparatus in the same time, P the height of the barometer, and p_s the pressure of saturated vapour as determined by Landolt, the densities obtained on the basis of Landolt's pressures, and given in the third column, are evidently represented by—

$$\frac{w(P - p_s)}{Wp}$$

The numbers of the fifth column, which are represented in the same way by—

$$\frac{w(P - p_R)}{Wp_s}$$

where p_R denotes the pressure as determined by Regnault's

* Liebig's Annalen, suppl. vi. (1868), p. 157.

† Chem. Abstr., vol. xiv., p. 736. The experiments date from 1869.

experiments, have been calculated by the present writer by multiplying the numbers of the third column by—

$$\frac{p_L (P - p_R)}{p_R (P - p_L)}$$

TABLE VII.—ACETIC ACID.

Determinations of Vapour-density by Distillation.

Temp.	Pressure according to Landolt.	Density observed, Horstmann & Landolt.	Pressure according to Regnault.	Density calc. from Regnault's pressures by eq. (12).	Density observed, Horstmann & Regnault.	I. Excess of observed density.	II.
25.0	23.5	2.42	15.13	3.86	3.80	-.06	
23.8	22.4	2.23	14.19	3.86	3.56		-.30
22.6	21.6	2.29	13.31	3.87	3.76	-.11	
21.5	20.4	2.24	12.54	3.87	3.68	-.19	
20.4	19.2	2.05	11.81	3.88	3.37		-.51
20.2	19.0	2.28	11.68	3.88	3.75	-.13	
20.0	18.9	2.13	11.56	3.88	3.52		-.36
17.4	16.8	2.09	9.95	3.89	3.56	-.33	
15.6	15.6	1.98	8.96	3.90	3.48	-.42	
15.3	15.3	1.95	8.81	3.90	3.42		-.48
15.3	15.3	1.85	8.81	3.90	3.24		-.66
14.7	15.1	1.78	8.54	3.91	3.18	-.73	
12.7	13.7	1.96	7.60	3.91	3.56	-.35	
12.4	13.5	1.89	7.46	3.92	3.45	-.47	

As the height of the barometer in Horstmann's experiments is not given it has been necessary to assume $P=760$. The inaccuracy due to this circumstance is evidently trifling. The last two columns of the table, which relate to different series of experiments by Horstmann (a distinction not observed in other parts of the table), give the excess of the densities thus obtained from Horstmann's and Regnault's experiments above the values calculated from equation (12) with the use of Regnault's determinations of pressure.

The densities obtained by experiment are without exception less than those obtained from equation (12). At the highest temperatures, where the liability to error is the least, both in respect to the measurement of the pressure of saturated vapour and in respect to the analysis of the product of distillation, the results of experiment are most uniform, and most nearly approach the numbers required by the formula. At the lowest temperatures the greatest observed density is about one-eleventh less than that required by the formula, the difference being about the same as between the highest and lowest observed values for the same temperature.

Since each successive purification of the substance employed by Regnault diminished the pressure of its vapour, it is not improbable that the pressures might have been still farther diminished by farther purification of the substance. The pressures which we have used are therefore liable to the suspicion of being too high, and it is quite possible that more accurate values of the pressure would still farther reduce the deficiency of observed density.

Perchloride of Phosphorus.—For this substance we have at atmospheric pressure a single determination of vapour-density by Mitscherlich,* and a series of determinations by Cahours;† at lower pressures we have determinations by Wurtz‡ and by Troost and Hautefeuille.§ In the experiments of Wurtz the pressure was reduced by mixing the vapour with air. In Table VIII. all these determinations are compared with the formula—

$$\log \frac{3.6(D-3.6)}{(7.2-D)^2} = \frac{544x}{t + 273} + \log p - 14.353. \quad (13)$$

The differences between the calculated and observed values are often large, in six cases exceeding 0.30; but

they exhibit in general that irregularity which is characteristic of errors of observation. We should expect large errors in the observed densities on account of the difficulty of obtaining the substance in a state of purity, and because the large value of the density renders it very sensitive to the effect of impurities which diminish the density,—also because the specific heat of the vapour is great, as shown by the numerator of the fraction in the second member of (13),* and because the density varies very rapidly with the temperature as seen by the numbers in the third column of Table VIII.

TABLE VIII.—PERCHLORIDE OF PHOSPHORUS.

Experiments of Mitscherlich, Cahours, Wurtz, and Troost and Hautefeuille.

Temperature.	Pressure.	Density calc. by eq. (13).	Density observed. Mitsch. Cahours.	Excess of observed density. Mitsch. Cahours.
336	(760)	3.610	3.656	+0.046
327	754	3.614	3.656	+0.042
300	765	3.637	3.654	+0.017
289	(760)	3.656	3.69	+0.034
288	763	3.659	3.67	+0.011
274	755	3.701	3.84	+0.139
250	751	3.862	3.991	+0.129
230	746	4.159	4.302	+0.142
222	753	4.344	4.85	+0.506
208	(760)	4.752	4.73	-.021
200	758	5.018	4.851	-.167
190	758	5.368	4.987	-.381
182	757	5.646	5.078	-.568
Wurtz. T. & H. Wurtz. T. & H.				
178.5	227.2	5.053	5.150	+0.097
175.8	253.7	5.223	5.235	+0.012
167.6	221.8	5.456	5.415	-.041
154.7	221	5.926	5.619	-.307
150.1	225	6.086	5.886	-.200
148.6	244	6.169	5.964	-.205
145	391	6.45	6.55	+0.10
145	311	6.37	6.70	+0.33
145	307	6.36	6.33	-.03
144.7	247	6.287	6.14	-.147
137	281	6.53	6.48	-.05
137	269	6.51	6.54	+0.03
137	243	6.48	6.46	-.02
137	234	6.47	6.42	-.05
137	148	6.31	6.47	+0.16
129	191	6.59	6.18	-.41
129	170	6.56	6.63	+0.07
129	165	6.55	6.31	-.24

But at the two lowest temperatures of Cahours's experiments the differences of the observed and calculated densities (0.381 and 0.568) are not only great, but exhibit, in connection with the adjacent numbers, a regularity which suggests a very different law from that of the formula. In fact, the densities obtained by Cahours at atmospheric pressure and those obtained by Troost and Hautefeuille at pressures a little less than one-third of an atmosphere seem to form a continuous series, notwithstanding the abrupt change of pressure. Yet it is difficult to admit that the density is independent of the pressure. So radical a difference between the behaviour of this substance and that of the others which we have been considering requires unequivocal evidence. Now it is worthy of notice that the experiment at 183°, in which the greatest discrepancy is seen, is not given in the first record of the experiments, which was in the *Comptes Rendus* in 1845. It is given in the *Annales de Chimie et de Physique* in 1847, where it is called the first experiment. (The experiment at 336° is also omitted in the *Comptes Rendus*, and that at 208° in the *Annales*; otherwise the lists are the same.) If it was the first experiment in point of time, which is apparently the meaning, it was made before the publication in the *Comptes*

* Pogg. Ann., vol. xxix. (1833), p. 221.

† Com. des Rendus, xxi. (1845), p. 625; and Ann. de Chim. et de Phys.,

3. vol. xx. (1847), p. 369.

‡ Comptes Rendus, vol. lxvii. (1878), p. 602.

§ Ibid., vol. lxxiii. (1876), p. 977.

* Compare Trans. Conn. Acad., vol. iii., p. 249.

Rendus, and we can only account for its omission by supposing that it was a preliminary experiment, in which its distinguished author did not feel sufficient confidence to include it at first with his other determinations, although he afterwards concluded to insert it. If we reject this observation as doubtful, the disagreement between the formula and observation appears to be within the limits of possible error, but additional experiments will be necessary to confirm the formula.*

Experiments have also been made by M. Wurtz, in which the vapour of the perchloride of phosphorus was diluted with that of the perchloride.† These experiments may be used to test equation (8), which, when the values of its constants are determined by equation (13), reduces to the form—

$$\log \frac{P_3}{P_2 P_1} = \frac{5441}{t_c + 273} - 13.751, \quad (14)$$

where P_1 , P_2 , and P_3 denote the partial pressures due respectively to the PCl_5 , the Cl_2 , and the PCl_3 existing as such in the gas-mixture. Since these quantities cannot be the subjects of immediate observation, a farther transformation of the equation would be convenient. Let M_2 , M_3 denote the quantities of the protochloride and of chlorine of which the mixture may be formed, and P_2 , P_3 the pressure which would belong to each of these if existing by itself with the same volume and temperature. These quantities will be connected with the equations—

$$P_2 = \frac{k t M_2}{2.22 v}, \quad P_3 = \frac{k t M_3}{4.98 v}, \quad (15)$$

where k denotes the same constant as before referred to. From the evident relations—

$$P_2 = p_2 + p_3, \quad P_3 = p_3 + p_5, \quad p = p_2 + p_3 + p_5,$$

we obtain—

$$p_3 = P_2 + P_3 - p, \quad p_2 = p - P_3, \quad p_5 = p - P_2;$$

and by substitution of these values in equation (14),—

$$\log \frac{P_2 + P_3 - p}{(p - P_2)(p - P_3)} = \frac{5441}{t_c + 273} - 13.751 \quad (16)$$

In view of the relations (15) this may be regarded as an equation between the pressure, the temperature, the volume, and the quantities of protochloride of phosphorus and chlorine in which the gas-mixture is resolvable.

It is in this form that we shall apply the equation to the experiments of M. Wurtz, the results of which are exhibited in Table IX. The first column gives the number distinguishing each experiment in the original memoir; the second, the temperature; the third, the observed pressure (p) of the mixture of PCl_5 , PCl_3 , and Cl_2 , which is the barometric pressure corrected for the small quantity of air remaining in the flask; the fourth, the pressure π due to the possible perchloride, found by subtracting the pressure due to the excess of protochloride (this pressure is calculated from the theoretical density of the protochloride) from the total pressure; the fifth, the density δ of the possible perchloride calculated from its pressure π with the temperature and volume. The numbers of these five columns are taken from the memoir cited, except that the correction of the barometric pressures has been applied by the present writer in accordance with the data furnished in that memoir. The two next columns contain the values of P_2 and P_3 . These would naturally be calculated from M_2 and M_3 by equations (15). But since the values of M_2 and M_3 have not been given explicitly, those of P_2 and

P_3 have been calculated from the recorded values of π and δ . Since the weight of the possible chloride is—

$$\frac{7.2}{2.22} M_2,$$

we have—

$$\delta = \frac{7.2 M_2 k t}{2.22 v \pi} = \frac{7.2}{\pi} P_2,$$

Moreover,—

$$p - \pi = P_3 - P_2,$$

since both members of the equation express the pressure due to the excess of the protochloride. The values of P_2 and P_3 were obtained by these equations.

No. of expt.	t_c	δ (obs.)	π	δ	P_2	P_3	p calc. by eq. (16).	Excess of obs. value of p .
XII.	173.29	7.61	423	6.68	392.4	725.5	760.7	-4.6
X.	165.4	748.4	413	6.80	390.1	725.3	747.9	+0.5
VII.	176.24	751.0	411	6.88	392.7	732.7	773.1	-22.1
VIII.	169.35	724.1	394	7.16	391.8	721.9	750.5	-26.4
V.	175.26	743.3	343	7.03	334.9	735.2	764.4	-21.1
II.	164.9	758.5	338	7.38	346.4	766.9	782.9	-24.4
XI.	175.75	760.0	318	7.00	303.2	751.2	776.8	-16.8
IV.	175.26	756.3	271	7.06	365.7	751.0	779.9	-14.6
IX.	160.47	753.5	214	7.44	221.1	760.6	766.8	-13.3
I.	165.4	760.0	194	7.25	195.3	761.3	768.5	-8.5
VI.	170.34	751.2	174	8.30	2.06	777.8	787.6	-36.4
III.	174.28	742.7	168	7.74	180.6	755.3	766.5	-23.8

The eighth column of the table gives the values of p calculated from the preceding values of t_c , P_2 , and P_3 , by equation (16); and the last column the difference of the observed and calculated values of p . The average difference is 18 mm., or a little more than 2 per cent, the observed pressure being almost uniformly less than the calculated value. This deficiency of pressure is doubtless to be accounted for by a fact which MM. Troost and Hautefeuille have noticed in this connection. The protochloride of phosphorus deviates quite appreciably from the laws of Mariotte, Gay-Lussac, and Avogadro, the product of the volume and pressure of a given quantity of vapour at 180° and the pressure of one atmosphere being 1548 per cent less than at the same temperature and the pressure of one-half an atmosphere.* Now we may assume, as a general rule, that when the product of volume and pressure of a gas is slightly less than the theoretical number (calculated by the laws of Mariotte, Gay-Lussac, and Avogadro) the difference for any same temperature is nearly proportional to the pressure.† It is therefore probable that between 160° and 180°, at pressures of about one atmosphere, the product of volume and pressure for protochloride of phosphorus is somewhat more than 3 per cent less than the theoretical number. The experiments of Wurtz, as exhibited in Table IX., show that the pressure, and therefore the product of volume and pressure (we may evidently give the volume any constant value as unity), in a mixture consisting principally of the protochloride, is on the average a little more than 2 per cent less than is demanded by theory, the differences being greater when the proportion of the protochloride is greater. The deviation from the calculated values is therefore in the same direction, and about such in quantity, as we should expect.‡

* Troost and Hautefeuille, *Comptes Rendus*, vol. lxxiii. (1876), p. 334.

† Andrews, "On the Gaseous State of Matter," *Phil. Trans.*, vol. clxvi. (1876), p. 447.

‡ The deviation of the protochloride of phosphorus from the laws of ideal gases shows the impossibility of any very close agreement between such equations as have been deduced in this paper and the results of experiment in the case of gas-mixtures in which the substance is one of the components. With respect to the question whether future experiments on the vapour of the perchloride alone, or with an excess of chlorine or of the protochloride, will reduce the disagreement between the calculated and observed values to such magnitudes as occur in the case of the protochloride alone, it would be rash to attempt to anticipate the result of experiments.

* Additional experiments on the density of this vapour have been made by M. C. Boura, concerning which he says in 1866: "Les déterminations qui je viens d'effectuer à 170 et 172 deg. (ce corps bout vers 160 à 165 degrés) m'ont donné des nombres qui, bien que notablement plus forts que ceux que j'ai obtenus antérieurement à 185 et 185 deg. ne sont pas bien éloignés de celui que correspond à 4 volumes." (*Comp. Rendus*, 63, p. 16.) So far as the present writer has been able to ascertain, these determinations have not been published. The formula gives 8.025 for 170° and 5.973 for 172°, at atmospheric pressure. The number corresponding to four volumes is 7.20.

† *Comptes Rendus*, vol. lxxvi. (1878), p. 601.

M. Wurtz has remarked that the average value of δ (the density of the possible perchlorides) is nearly identical with the theoretical density of the perchloride, and appears inclined to attribute the variations from this value to the errors of experiment. Yet it appears very distinctly in Table IX., in which the experiments are arranged according to the value of π (the pressure due to the possible perchloride), that δ increases as π diminishes. The experiments of M.M. Troost and Hauteville show that the coincidence remarked by M. Wurtz is due to the fact that on the average in these experiments the deficiency of the density of the possible perchloride (compared with the theoretical value) is counterbalanced by the excess of density of the protochloride. When $\pi > 400$, the effect of the deficiency in the density of the possible perchloride distinctly preponderates; when $\pi < 250$, the effect of the excess of density in the protochloride distinctly preponderates. But the magnitude of the differences concerned is not such as to invalidate the general conclusion established by the experiments of M. Wurtz, that the dissociation of the perchloride may be prevented (at least approximately) by mixing it with a large quantity of the protochloride.

TABLE X.

For the solution of the equation: $\log \frac{1000 (\Delta - 1)}{(2 - \Delta)} = L$.

L	Δ	Diff.	L	Δ	Diff.
0.7	1.005	1	4.2	1.778	22
0.8	1.006	2	4.3	1.800	19
0.9	1.008	2	4.4	1.819	18
1.0	1.010	2	4.5	1.837	17
1.1	1.012	2	4.6	1.854	14
1.2	1.015	3	4.7	1.868	14
1.3	1.019	4	4.8	1.882	12
1.4	1.024	5	4.9	1.894	11
1.5	1.030	6	5.0	1.905	10
1.6	1.037	7	5.1	1.915	9
1.7	1.046	9	5.2	1.924	8
1.8	1.056	10	5.3	1.932	7
1.9	1.069	13	5.4	1.939	6
2.0	1.084	15	5.5	1.945	6
2.1	1.102	18	5.6	1.951	5
2.2	1.122	20	5.7	1.956	5
2.3	1.146	24	5.8	1.961	5
2.4	1.172	26	5.9	1.965	4
2.5	1.202	30	6.0	1.969	4
2.6	1.234	32	6.1	1.972	3
2.7	1.268	34	6.2	1.975	3
2.8	1.305	37	6.3	1.978	3
2.9	1.343	38	6.4	1.980	2
3.0	1.382	39	6.5	1.982	2
3.1	1.421	39	6.6	1.984	2
3.2	1.461	40	6.7	1.986	1
3.3	1.500	39	6.8	1.987	1
3.4	1.537	37	6.9	1.989	1
3.5	1.574	35	7.0	1.990	
3.6	1.609	33	7.2	1.992	
3.7	1.642	31	7.4	1.994	
3.8	1.673	30	7.6	1.995	
3.9	1.703	27	7.8	1.996	
4.0	1.730	25	8.0	1.997	
4.1	1.755	23	9.0	1.999	
4.2	1.778				

Table for facilitating calculation.—The numerical solution of equations (10), (11), (12), and (13), for given values of t and p , may be facilitated by the use of a table, if we set—

$$\Delta = \frac{D}{D_1} \quad (17)$$

$$L = \log \frac{1000 D_1 (D - D_1)}{2 D_1 - D} = \log \frac{1000 (\Delta - 1)}{(2 - \Delta)} \quad (18)$$

we have for peroxide of nitrogen,—

$$L = \frac{3.1586}{tc + 273} + \log p - 9.451 \quad (19)$$

for formic acid,—

$$L = \frac{3800}{tc + 273} + \log p - 9.641 \quad (20)$$

for acetic acid,—

$$L = \frac{3520}{tc + 273} + \log p - 8.349 \quad (21)$$

and for perchloride of phosphorus,—

$$L = \frac{5441}{tc + 273} + \log p - 11.353 \quad (22)$$

By these equations the values of L are easily calculated. The values of Δ may then be obtained by inspection (with interpolation when necessary) of the following table. From Δ the value of D may be obtained by multiplying by D_1 , viz., by 1.589 for peroxide of nitrogen or formic acid, by 2.073 for acetic acid, and by 3.6 for perchloride of phosphorus.*

The constants of these equations are of course subject to corrections by future experiments, which must also decide the more general question—in what cases, and within what limits, and with what degree of approximation, the actual relations can be expressed by equations of such form. In the case of perchloride of phosphorus especially the formula proposed requires confirmation.—*American Journal of Science and Arts.*

PROCEEDINGS OF SOCIETIES

PHILOSOPHICAL SOCIETY OF GLASGOW. CHEMICAL SECTION.

Mr. JAMES MACTEAR, President, in the Chair.

At the Annual Business Meeting of this Society on November 24, a short paper was read by Mr. J. B. HANNAY on "Crystalline Cavities," in which he brought forward some theories as to the possible formation of crystals.

The President's Address was read at the next meeting, December 8, an abstract of which is given below.

"On the Antiquity of Chemical Science," by JAMES MACTEAR, F.C.S., F.I.C., Member of the International Jury, Paris, 1878.

Taking up the subject at the point where the science is usually supposed to have originated—in the Arabian School of Philosophy in the 8th century—the author showed that the science was of much more ancient origin, and traced its history back through the medical science of the Greeks, Russians, Egyptians, &c., to Indian sources, where in all probability it was known and cultivated as a science fully 2000 B.C. The author also showed the unsatisfactory nature of the derivation of the name of the science from Chem or Chema, as meaning dark, hidden, or black, or even Dshem, to find, as it has been explained by all previous authors on the subject, and stated that his own researches into the subject resulted in deriving the name indirectly from the word Knams—an Arabic word of Sanskrit origin—meaning fire, or rather from its compound Khamis, meaning the fifth. As is well known the ancient Hindoos recognized five elementary types—water, fire, earth, air, and ether, the latter being the purest.

* The value of Δ diminished by unity expresses the ratio of the number of the molecules of the more complex type to the whole number of molecules. Thus, if $\Delta = 1.70$, in the case of peroxide of nitrogen there are 20 molecules of the type N_2O_4 to 30 of the type N_2O_2 , or in the case of perchloride of phosphorus there are 20 molecules of the type PCl_5 to 40 of the type PCl_3 and 40 of the type P_2Cl_4 . A consideration of the varying values of Δ is therefore more instructive than that of the values of D , and it would in some respects be better to make the comparison of theory and experiment with respect to the values of Δ .

ciple, or type, of active force or motion which caused the changes in the condition of the elementary types or their combinations. From a consideration of those and other facts the author derives the title of the science from—

AL-KHAMIS,
(THE FIFTH),

meaning *The Science of Force or Change*. No more perfect descriptive title could be found for it, even in our present enlightened age.

A cordial vote of thanks was given to Mr. Maclear for his interesting paper.

NOTICES OF BOOKS.

Technical Vocabulary, English and German. Technisches Vocabular für Technische Lehranstalten und zum Selbststudium für Studierende. Techniker und Industrielle. Von Dr. F. J. WERSHOVEN, mit einem Vorwort von A. VON KAVEN, Direktor der Technischen Hochschule in Aachen. Leipzig: Brockhaus.

As it is intimated in the preface, the student, in acquiring a foreign language, learns every portion of it sooner than the technical phraseology of the sciences and the useful arts. The ordinary grammars, vocabularies, and dictionaries avoid such subjects,—often, indeed, from ignorance on the part of their authors. That such a state of things should be perpetuated in an eminently industrial age is no less strange than lamentable. To an Englishman, surely the French or the German of the laboratory and the workshop must be more important than the French or German of the feuilletonist and the play-wright. We must therefore welcome every attempt to render the technological literature of each country more widely intelligible among its neighbours.

The present little work may be pronounced remarkable for its general accuracy. We find, under the alkali manufacture, "boot-pan" in place of "boat-pan," but this is a mere misprint. The terms "metalloids" for non-metallic bodies, and "Daltonism" for colour-blindness, are now rarely, if at all, used by accurate English writers.

As a matter of course, when a writer gives in the compass of 220 pages the technical phraseology of mechanics, of the various branches of physics, of chemistry, meteorology, geology, metallurgy, photography, galvanoplastics, railways, land surveying, glass-making, pottery, textile manufactures, paper-making, sugar-extraction, and brewing, he cannot be expected to go into every detail in any of these sciences and arts. But no one, we submit, could have done more in a space so limited. Some of the most important chemical arts have certainly been omitted, such as dyeing, tissue-printing, colour-making, tanning, soap-making, &c. Perhaps it might have been better if the author had divided the matter into two distinct volumes, a mechanical and a chemical, since many students and technologists who are deeply interested in the arts just enumerated will find certain sections of this work, e.g., that on railways, quite out of their sphere, and *vice versa*.

Still, whilst we venture to throw out these suggestions, we can do no other than pronounce the work, such as it is, a true and an accurate *multum in parvo*.

Royal Institution.—The Managers have awarded the Addison Prize of £105 to G. S. Boutger, F.L.S., F.G.S., for an essay on "The Structure and Functions of the Retina in all Classes of Animals, viewed in Relation with the Theory of Evolution." Professor Tyr. all will give the first of his Christmas Lectures (adapted to an educated juvenile auditory) on Saturday, December 27, at 3 o'clock.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Comptes Rendus Hebdomadaires des Séances, l'Académie des Sciences. No 22, December 1, 1879.

Observations on M. Trécul's last notice on Chlorophyll.—M. Chevreul.—The author asks, first, Whether M. Trécul has found the crystalline form of chlorophyll soluble without residue in alcohol and ether? This question, having been answered in the affirmative, he expresses his regret at not having known of his discovery in 1865. He then asks whether M. Trécul regards chlorophyll as a simple product of the organism, as would appear from its solubility in alcohol and ether and from its crystalline character? M. Trécul's opinion is that the chlorophyll which he terms globular is a living organ, the producer of the chemical principle crystalline chlorophyll, and the reducer of carbonic acid in the green living leaf. M. Chevreul then, referring to certain interesting experiments by M. Croëz, asks, What is the part played by chlorophyll? Is it a constituent part of the organ, or its presence merely accessory?

Certain Properties of Glucose.—E. Peligot.—On treating glucose with alkalis, the author has obtained a crystalline compound to which he gives the name of saccharin. Its composition, $C_{12}H_{11}O_{11}$, is that of saccharose, but it does not ferment in presence of yeast, and it is almost insipid, with a faint after taste recalling that of Glauber's salt. It is only reduced by the cupric alkalis after a prolonged boiling.

Crystalline Form and Optical Properties of Saccharin.—M. des Cloizeaux.

Measurement of Quantities of Electricity.—G. A. Hirn.—The author criticises the various methods proposed for measuring electricity by its effect, and in particular to the recent investigations of Prof. Villari.

Remarkable Electric Phenomenon.—E. Lamarre.—At the commencement of a violent snowstorm the author perceived little luminous tufts at the end of each of the steel rods of his umbrella, and heard a sound like the humming of an insect. On approaching his hand to one of the lights he felt a slight shock in the first two joints, and the light disappeared.

Separation of Phosphoric Acid from the Sesquioxides of Iron and Alumina.—P. Derome.—See p. 292.

Composition of the Horn of the Stag.—A. Bleunard.—Stag's horn is a lower homologue of coagulated egg-albumen, but is more highly hydrated.

Determination of Chlorine in Various Seeds and Forage Plants.—R. Nöke.—The author gives a table of his results, which differ from the older determinations, and show that chlorides form a part of every vegetable diet.

Berichte der Deutschen Chemischen Gesellschaft zu Berlin, No. 13, 1879.

Formula of Quinhydrone.—H. Wichelhaus.—The author considers that the formula $C_{18}H_{14}O_6$ is established beyond doubt.

Reaction for Phenyl-glyoxylic Acid.—L. Claisen.—If to a solution of this acid in benzol there is added concentrated sulphuric acid the mixture, after standing for a short time and agitation, takes a deep red colour, which soon passes into a violet-blue. On the addition of water the colouring-matter passes with an intense crimson hue into the supernatant layer of benzol, from which it may be separated by evaporation, or on the addition of petroleum-ether. The derivatives of the acid, its amides and ethers, and benzoyl cyanide behave in a similar

manner. Meta-nitro-phenyl-glyoxylic acid yields a splendid crimson-red colouration and ortho-nitro-benzoyl-cyanide a blue-green.

Aldehyd Collidin.—A. Wischnegradsky.—On oxidising collidin by means of chromic acid the author obtained an acid, $C_8H_7NO_4$, forming fine white prisms, readily soluble in hot water, and easily volatilised by heat, though with partial carbonisation.

Decomposition of Ethylamin Hydrochlorate by the Action of Heat.—M. Fileti and A. Piccini.—The gases evolved consist of a mixture of ammonia, mono- and di-ethylamin, ethyl-chloride, and ethylen.

A New Method of the Formation of Hyponitrous Acid and of Hydroxylamin.—W. Zorn.—If in the electrolysis of sodium nitrite a mercury electrode is employed at the negative pole, and the process is interrupted when all the nitrite is consumed, then, on adding acetic acid and silver nitrate, nitro-silver is deposited in abundance. Hydroxylamin is also produced.

Behaviour of Cymol in the Animal Body.—Oscar Jacobsen.—Cuminoic acid, $C_{12}H_{15}NO_3$ appears in the urine. The author describes its properties, and those of certain of its salts.

A Remarkable Oxidation-product of Cholic Acid.—P. Latschinoff.—On oxidising impure cholic acid with potassium permanganate or nitric acid, the author obtains a mixture of crystallised acids, of which he here describes one, the choloidanic acid. No fixed fatty acids were obtained on a similar treatment of pure cholic acid. Choloidanic acid is isomeric with camphoric acid, but distinct from all the isomers of the latter hitherto described.

Direct Separation of Manganese and Iron.—F. Beilstein and L. Jawein.—The authors propose two processes. In the one the solution of the two metals is poured into an excess of a concentrated aqueous solution of potassium cyanide. After standing from half an hour to an hour the precipitate re-dissolves, and only a slight turbidity remains. The whole is filtered, the residue is dissolved in a few drops of dilute hydrochloric acid, the solution is mixed with excess of potassium cyanide, and the clear solution is added to the filtrate. All these operations are performed in the cold. Solid iodine is then added till the liquid appears brown, when any traces of free iodine are removed by the addition of a few drops of free alkali. The complete separation of the manganese is ascertained by adding iodine to a small portion of the liquid, decanted or filtered; heating very gently and adding soda-lye. The liquid should remain clear. The precipitated manganic oxide is filtered off, washed, dissolved in hydrochloric acid, precipitated with ammonium sulphide at a boil, and weighed as manganese sulphide. Second process:—The salt of the two metals is dissolved in concentrated nitric acid (1.35), heated to a boil and kept at this temperature, whilst potassium chlorate is gradually added in small portions. The liquid is diluted with water and filtered. The precipitate when washed is always found to contain iron. It is therefore re-dissolved in hydrochloric acid, evaporated to dryness, and the residue taken up in concentrated nitric acid, and treated again with chlorate of potassa at a boil. A peroxide is now precipitated containing very small traces of iron.

Determination of Acetyl by Means of Magnesia.—Hugo Schiff.—The magnesia used must be precipitated from magnesium sulphate or chloride (free from iron), avoiding excess of alkali, and kept as a paste under water.

Constitution of Ellagic Acid.—Hugo Schiff.—Not suitable for abstraction.

Contributions to a Knowledge of the Spontaneous Oxidation of Manganese Monoxide, with Reference to the Regeneration of Manganese.—J. Post.—The rapidity of the action is much promoted by exposing manganese oxide to the air in a thin layer. On the small scale an addition of soap with the aid of constant agita-

tion was found advantageous. The addition of an excess of alkali or of lime increases the yield, the former being more beneficial.

Hydroxy-isobutyl-formic Acid.—W. von Miller.—The compound formed on the oxidation of isobutyl-formic acid by potassium permanganate, and which precedes the formation of dimethyl-acrylic acid, is β -oxy-isobutyl-formic acid.

Hydroxy-ethyl-methyl-acetic Acid.—W. von Miller.—On distilling this acid with sulphuric acid there was formed no methyl-crotonic acid.

A New Salt of an Iridium Base.—K. Birnbaum.—The author points out that most of the researches on the iridium bases date from a time when the metal had not been obtained in a state of purity.

Behaviour of Anhydrous Calcium Oxide with Anhydrous Carbonic Acid.—K. Birnbaum and M. Mahn.—The melting-point of zinc is the temperature at which anhydrous calcic oxide begins to absorb carbonic acid under the ordinary atmospheric pressure. A definite compound is not formed, as at this temperature calcium carbonate is within the limits of dissociation.

Excitement of Oxygen by Nascent Hydrogen.—F. Hoppe-Sevler.—The author considers that every attempt to explain the vital process in plants and animals necessarily demands the assumption of a plentiful source of the activation of oxygen within the organism. He has convinced himself that the formation of free hydrogen takes place only in the absence of oxygen, and that on the access of oxygen to putrescent liquids not alone the nascent hydrogen is oxidised, but that energetic processes of oxidation are set up.

On Chlorophyll.—F. Hoppe-Sevler.—The author is engaged with the examination of chlorophyllan, a crystalline colouring-matter, obtained from the residue of the alcoholic extract of grass, after the erythrophyll of Boegarell has crystallised out.

A Correction.—W. Ramsay.—Refers to *Berichte*, xii., p. 1024.

An Explanation.—C. Willgerodt.—See *Berichte*, xi., 1792, and xii., 1319.

Perchloric Acid as a Reagent for Alkaloids.—G. Fraude.—The author has examined the action of perchloric acid upon the alkaloids of the quinin group, the opium bases, veratrin, caffeine, atropin, nicotin, and coniin. All these give no colour-reactions, which, however, occur with aspidospermin and the strychnos bases. The colourations are sufficiently permanent to admit of spectroscopic examination.

On Aspidospermin.—G. Fraude.—There are two varieties of aspidospermin obtained respectively from two varieties of the *Aspidosperma quebracho*, the "blanco," and the "colorado." The author gives an account of some of the salts of aspidospermin and its reactions.

Maleic Acid from Dichloroacetic Acid.—S. Tannar.—The nature of this paper will appear from the title.

Amido-anthraquinon from Anthraquinon-mono-sulphonic Acid.—H. R. v. Perger.—The author has examined the action of ammonia upon the above-mentioned acid.

Action of Phosphorus Penta-chloride and Hydriodic Acid upon Saccharic Acid.—H. de la Motte.—The author, having heated pure potassium saccharate with 6 molecules of phosphorus penta-chloride, obtained chloromuconic acid, $C_6H_4Cl_2O_4$. He finds that this acid, whether obtained from saccharic acid or mucic acid, contains, further, 2 mols. crystalline water.

Certain Derivatives of Santonin. S. Cannizzaro and G. Carnelutti.—Santonin, mixed with red phosphorus and heated for a long time to ebullition with an excess of concentrated hydriodic acid, yields santonos acid, a powerful monobasic compound, containing an atom of oxygen

ess than the isomeric santonin acid. The authors have examined the methyl-ether of this acid.

The so-called Digallic Acid.—P. Freda.—The substance characterised by Schiff as digallic acid cannot be obtained.

Action of Boron Fluoride upon Aceton.—F. Landolph.—The author has obtained and isolated α -fluoboric aceton and a β -fluoboric aceton.

Two New Hydrofluoboric Acids and Fluoboric Ethylen.—F. Landolph.—The respective formulæ of these acids are $\text{Bo}_2\text{O}_7\text{H}_4\cdot 3\text{HFl}$ and $\text{Bo}_2\text{O}_5\text{H}_4\cdot 2\text{HFl}$.

Analysis of Organic Fluor and Boro Compounds.—F. Landolph.—Not suitable for abstraction.

Potassa-melt of Rhamnetin.—St. Smorawski.—Among the products obtained are proto-catechuic acid and phloro-glucin. A substance which, like the quercetic acid of Hlasiwetz, gives with alkalis an intense red reaction accompanied the crude proto-catechuic acid.

Sequel to the Decomposition of the Oxy-anthraquinones by Potassa.—C. L. ebermann and J. Dehnst.—Para-oxy-benzoic acid was observed along with benzoic and proto-catechuic acid in the potassa melt of mono-sulpho-anthraquinonic acid.

Constitution of the Camphor Compounds.—M. Balle.—Not suitable for abstraction.

Behaviour of Indigo-white with Potassium Pyrosulphate.—A. Baeyer.—There exists a compound of indigo-white which possesses the general properties of Baumann's potassium sulph-indoxylate, and according to its origin is a sulpho-phenolate.

Aromatic Sulpho-ureas.—C. Feuerlein.—The author has analysed the phenyl-cyan-imid hydrate, and has succeeded in forming a platinum and a silver compound. The attempts at preparing the platinum and the silver salts of mono-phenyl-guanidin gave only very small precipitates, which proved to be the corresponding phenyl-cyanimide compounds.

The Acid of Drosera Intermedia.—G. Stein.—The acid in question is the citric.

Changes of Ammonium Isæthionate at Elevated Temperatures.—F. Carl.—The results indicated that the substance melting at 196° to 198° is the ammonium salt of a new acid, the di-isæthionic, formed by the union of 2 mols. ammonium isæthionate, with the simultaneous removal of a mol. of water.

The Constitution of the Phenyl Halogen Propionic Acids.—E. Erlenmeyer.—Not suitable for abstraction.

Certain New Colouring-matters.—P. Greiff.—If one part of chloranil is allowed to act upon two parts of dimethyl-anilin a deep blue colouration is observed, even in the cold. On heating to 50° the reaction is complete, and a bronze-coloured melt is obtained, which is insoluble in water, but gives up to alcohol and acetic acid a deep violet-blue colour of great purity. If instead of dimethyl-anilin, methyl-diphenyl-amin is employed, a blue colour is obtained, more beautiful than any yet known. The reactions are very complete and the yield is high.

New Determinations of the Densities of Solid Organic Compounds.—H. Schroeder.—This important paper consists chiefly of tabulated results. The author remarks that there are 29 compounds, 23 of which contain the benzol-nucleus, whose volumes are almost exact multiples of 5.91. The elements carbon, hydrogen, oxygen, and nitrogen in the solid state generally take up the space of 1 stere.

Oxidation of Cholic Acid.—H. Tappeiner.—A controversial notice, having reference to the paper of Latschinoff (*Berichte*, xii., 1022).

Diffusion-Experiments on Solutions of Saline Mixtures of an Acid Reaction.—F. Hinteregger.—In most cases the acid diffuses more readily than the base, deviations from this rule occur, as in the case of phos-

phates. The mixture of free hippuric acid and sodium hippurate behaves in the very opposite manner.

Quinon-like Body Occurring in *Ag. atrotomentosus*.—W. Thörner.—Not susceptible of useful abstraction.

A New Organic Acid in *Agaricus Integer*.—W. Thörner.—The composition of this acid is $\text{C}_{15}\text{H}_{30}\text{O}_2$. It crystallises from alcohol in snow-white needles, arranged in little tufts and melting at $69\frac{1}{2}^\circ$ to 70° .

Action of Hydrocyanic Acid upon certain Diazo Compounds.—S. Gabriel.—It has hitherto not been shown that the cyanogen residue can be simply transferred by the action of compounds containing the cyanogen group upon the diazo compounds.

Certain Derivatives of Sulphacetic Acid.—S. Gabriel.—The author has succeeded in producing acids of the formula $\text{R}_{11}(\text{S}\cdot\text{CH}_2\cdot\text{COOH})_2$ from bibasic mercaptans and phenyls.

Action of Ammonia and the Amines upon the Quinons.—T. Zincke.—In consequence of the notice issued by Sommaruga (xii., 979) with respect to the investigation of these reactions, the author publishes the results on the same subject previously obtained by some of his pupils.

The Sugar of Populin.—E. O. von Lippmann.—The sugar obtained on the decomposition of populin is glucose.

Occurrence of Tricarballic Acid and Aconitic Acid in Beet-juice.—E. O. von Lippmann.—The citric, aconitic, and tricarballic acid accompany sugar in beet-juice.

The Phthalein of Hæmatoxylin.—E. A. Letts.—The author has not succeeded in obtaining this phthalein in the form of crystals. The alcoholic solution dries up to a gummy mass insoluble in water. With potassa-lye the phthalein yields a purple-red colouration totally different from that which the same reagent gives with solution of hæmatoxylin.

Certain Derivatives of Tri-ethyl Citrate.—J. Conen.—The author describes the action of phosphorus trichloride and penta-chloride upon this ether.

Formation of Potassium Ferro-chromate and the New Corresponding Ammonium Compound.—C. Hensgen.—The contents of this paper will sufficiently appear from the title.

Behaviour of Ammoniac Gum Resin on Distillation with Zinc Powder.—G. L. Ciamician.—The products of this gum-resin (salicylic acid, ortho-ethyl-phenol, &c.) are very different from those obtained by a similar process from the so-called terpen-resins. The portions boiling at higher temperatures in the case of gum-ammoniac consist of one of the higher hydrocarbons of the benzol-series, whilst from abietic acid and gum-elemi we obtain hydrocarbons of the naphthalin series.

Lauric Acid and its Transformation into Undecylic Acid.—F. Krafft.

Tri-decylic Acid, Penta-decylic Acid, and Margaric Acid.—F. Krafft.—These two papers do not admit of useful abstraction.

Cynurenic Acid.—M. Kretschky.—On treatment with zinc powder in a current of hydrogen this acid yielded quinolin almost perfectly pure.

State of the Alkaline Phosphates in Aqueous Solution.—J. M. van Bemmelen.—The record of diffusion experiments. In every moment a certain number of molecules of Na_3PO_4 are formed and again decomposed, the number being a function of the temperature and the quantity of water.

The Existence of Double Salts in Solutions.—P. H. B. Ingenhous.—The author's experiments refer chiefly to barium aceto-nitrate, calcium aceto-chloride, and osmium formio-nitrate. The results are shown in the form of tables.

Condensation Products of Tertiary Aromatic Bases.—Otto Fischer.—The author's researches embrace the oil of bitter almonds and dimethyl-anilin, the hydrochlorate of tetra-methyl-diamido-triphenyl-methan, bitter almond oil green (benzoyl green), cuminol and dimethyl-anilin, methylal and dimethyl-anilin, benzo-phenon chloride and dimethyl-anilin, and the phthalein of dimethyl-anilin.

Condensation Products of the Aldehyds with Primary Aromatic Bases.—Otto Fischer.—The author has obtained a new base, to which he gives the name of diamido-triphenyl-methan.

Apparatus for Determining the Heat of Combustion.—F. Fischer.

Apparatus for the Determination of Oxygen in Atmospheric Air.—F. Fischer.—These two papers cannot be intelligibly abstracted without the accompanying illustrations.

La Correspondance Scientifique.
October 28, 1879.

Action of Solar Light upon Vegetation in Northern Regions.—Nicolas de Nasakine.—The author maintains that the aroma of fruits increases with the latitude, while the sweetness decreases. The foliage of northern trees is remarkable for its deep shades of green, and the flowers have colours exceptionally vivid. Many herbs, such as carraway, are richer in essential oils in Norway than in more southern regions. The author ascribes these differences to the influence of the prolonged light of the summer months.

MISCELLANEOUS.

Royal Academy of Arts.—At a general assembly of Academicians, held on the 5th instant, Mr. A. H. Church, late of Cirencester, was elected to the Professorship of Chemistry in the Royal Academy.

Separation of Fats and Soaps.—Justus Wolff.—The mixture of soap and fat is comminuted as far as possible, mixed with from 10 to 20 parts of purified aniline, and digested in the water-bath, stirring and crushing continually for half to three-quarters of an hour. The solution is filtered when cold, and the residue is several times treated with aniline. In the residue is found all the soap, whilst the resins and fats are in the solutions. These are poured together, mixed with hydrochloric acid and with 3 or 4 parts of water, and when cold shaken with ether. The ethereal solution on evaporation leaves the unsaponified fats and resins. The aniline must be free from benzol and nitro-benzol.—*Zeitsch. Anal. Chemie*, xviii., 570.

NOTES AND QUERIES.

Association of Sulphuric Acid Manufacturers.—Can any reader of the *CHEMICAL NEWS* furnish me with the name and address of the secretary or other official representative of an Association formed last year among sulphuric acid manufacturers, with the view of developing sources of supply of sulphur other than pyrites, in consequence of the combination among pyrites companies to raise the price of pyrites?—VITRIOL.

MEETINGS FOR THE WEEK

MONDAY, 22nd—London Institution, 5.
Society of Arts, 8. (Cantor Lecture). "Chemistry of Bread and Bread Making," Dr. Graham.
TUESDAY, 23rd—Civil Engineers, 8. (Anniversary).
WEDNESDAY, 24th—Royal Institution, 3. Prof Tyndall. First Christmas Lecture on "Water and Air."

NOTICE.

Friday next being a Bank Holiday, THE *CHEMICAL NEWS* will be published on Wednesday, the 24th inst. Advertisements should reach the Office before 2 p.m. on Tuesday, the 23rd inst.

Box Court, Ludgate Hill, December 18, 1879.

PROFESSOR TYNDALL'S CHRISTMAS LECTURES

AT THE
ROYAL INSTITUTION.
Subject:—**WATER AND AIR.**

These Lectures will, by permission of Dr. TYNDALL, be reported verbatim for THE *JOURNAL OF SCIENCE*. Lecture I. will appear in the January number.

Seventeenth Year of Publication.

THE
JOURNAL OF SCIENCE.
(MONTHLY, FORMERLY "THE QUARTERLY JOURNAL OF SCIENCE")
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UNIVERSITY COLLEGE, BRISTOL.

CHEMICAL PROFESSORSHIP.

The Council invite Applications for the Chair of Chemistry. Salary, £300, with a share of the Students' Fees. Applications, with testimonials, to be sent not later than 9th February, 1880. Further information may be obtained from the Principal on application to EDWARD STOCK, M.R.C.S., Secretary.

The Gas Committee of the Corporation of Manchester are prepared to receive Tenders for the supply of Oxide of Iron, suitable for purification purposes to their Glynor and Rochdale Road Works, in such quantities as may be from time to time required during twelve months, commencing next spring.

Forms of Tender and full particulars can be obtained on application to Mr. George B. Jackson, Town Hall, Manchester.

Tenders endorsed "Tender for Oxide of Iron" and addressed to the Chairman of the Gas Committee must be delivered here not later than Wednesday, the 14th day of January next. The Corporation do not bind themselves to accept the lowest or any tender. By order,

JOS. HERON, Town Clerk.

Town Hall, Manchester,
December 16, 1879.

The Prestoles Alkali Works, Farnworth, near Bolton, Lancashire, fitted with costly plant, machinery, and apparatus for the manufacture of soda-ash, bleaching-powder and liquor, and sulphuric acid, in complete working order. With possession

MESSRS. FULLER, HORSEY, SONS, AND CO. are instructed to SELL by AUCTION, at the Pavilion Hotel, Manchester, on Friday, January 16th, at 3 p.m. exactly, in one lot (unless an acceptable offer be previously made by private contract), the PRESTOLE ALKALI WORKS, a freehold property, having a superficial area of 95,741 sq. yards of land, subject to a rent amounting to £407 13s. 9d. per annum, with the buildings, plant, machinery, and apparatus erected thereon capable of manufacturing monthly a product exceeding in the aggregate 2000 tons of soda-ash, bleaching-powder and liquor (by Weldon's patent process), and sulphuric acid, also caustic soda and muriatic acid. The amount expended in the construction of these works has been very large. A valuation was made with great care in 1874 by Messrs. H. James and Son, the well-known valuers, of Manchester; their estimate then amounted to £99,000, and since that time a sum exceeding £10,000 has been expended. The whole of the works and plant have been well kept, and are conveniently arranged for working. The Bury, Bolton, and Manchester Canal, which forms one boundary of the property, affords a facility for economical water carriage, and the Lancashire and Yorkshire Railway is within a very short distance. There is a plentiful supply of water for all manufacturing purposes free of cost, from the river Croal, and coal is raised from pits in the immediate neighbourhood. Pyrites, salt, lime, and limestone are all brought by boats direct to the wharf on the canal. The works may be economically worked with a small capital, as arrangements may be made with the vendors for a very considerable portion of the purchase-money to remain upon mortgage. Large profits have been realised in the past, and the high reputation of the Prestole manufacturers in the trade will ensure to an energetic man, even at the existing low prices, a ample return for his capital invested, and possession of works not surpassed for completeness or completeness by any in the kingdom. May be viewed till the sale. Printed particulars may be had at the works; at the Pavilion and Queen's Hotel, Manchester; of Mr. W. W. Wadman and Black, Solicitors, Warmistree; of Messrs. Christopher and Son, Solicitors, 28, Abchurch Lane, London; of Messrs. Fuller, Horsey, Sons, and Co., 11, Mill Lane, Square, London, who are empowered to treat for the disposal by private contract, or as offer for renting would be entertained.

Water-Glass, or Soluble Silicates of Soda and Potash, in large or small quantities, and either solid or in solution, at ROBERT KUMNEY'S, A. & C. Chemical Works Manchester.

THE CHEMICAL NEWS.

VOL. XL. No. 1048.

THE LAW OF THE CHEMICAL ELEMENTS.

By D. MENDELEEF.

(Continued from page 292.)

II. ON THE APPLICATION OF THE PERIODIC LAW TO THE SYSTEM OF ELEMENTS.

THE system of elements possesses not only a purely pedagogic importance as a means of learning more easily the different facts which are grouped in a systematic manner and united one to another, but also a scientific importance, inasmuch that it shows us new analogies, and thus opens up new roads for the research for elements. All the systems known up to the present can be placed in two distinct categories.

Into one of these categories (artificial systems) enter the systems which are based on some few characters of the elements. For example, the systems of division of the elements according to their affinity, according to their electro-chemical properties, according to their physical properties (or division into metals and metalloids), according to the manner in which they behave in the presence of oxygen and of hydrogen, according to their atomicity, &c. These systems, in spite of their evident defects, deserve some consideration, for they have the merit of a certain exactness, and each of them has contributed in a special direction to the progressive elaboration of chemical ideas.

The systems of the second category (natural systems) establish, according to members of different and purely chemical properties, groups of analogous elements. The well-known results of these systems have obtained for them the preference over the artificial systems. However, they are also stained with serious faults.

1st. The partition of elements does not rest upon invariable principles; also such elements as Tl, and even Ag and Hg, &c., have been placed in different groups. Na and K, Li and Rb, and often Tl also, have been frequently placed among the alkaline metals, although there exist between K and Na, in spite of the small difference between their atomic weights, more difference of properties than there is between K, Rb, and Cs. Na and all compounds of Na have a greater density than K and its corresponding compounds, although the atoms of sodium are lighter than the atoms of potassium.

NaCl and KCl crystallise in cubes (KCl often crystallises in a mixture of cubes and octahedra). However, we do not meet with isomorphous mixtures in nature, and we have obtained none artificially. According to Stassfurt we often meet with distinctly separate crystals of NaCl and KCl, one by the side of the other.

In the same manner the situation of Mg relative to Ca, or of Pb relative to Ba, Sr, and Ca, or of Tl relative to K, Rb, and Cs, is doubtful. In consequence of this want of fixed principles of separation the results of the natural systems are very uncertain.

2ndly. Some elements are without analogies, such as Au, Al, B, F, Ur, &c.

3rdly. There is no general expression for the reciprocal relations between the different groups; they could not be united in one whole. Thus these systems are always incomplete.

The periodic law possesses, in the forms of oxides and in the atomic weights, invariable numerical values for the distribution of the elements. It permits us, then, to group together the elements which have in reality a great deal

of resemblance, and it answers at the same time to the principles which have successively led to artificial systems. It gives us the means to erect an arbitrary system, as complete as possible. The preceding considerations, as well as those which are to follow, show up its advantages and properties. I will only insist on the manner of employing it to determine the places of some elements which have given rise to different interpretations. I must now commence with some general remarks.

The position of an element, R, in the system is found by the series and the group to which this element belongs, and therefore by the elements X and Y, which are next to it, one in front and the other behind in the same series, and also by two elements in the same group, the element R', whose atomic weight is the next smaller, and the element R'', whose atomic weight is the next larger. The properties of R can be determined from the known properties of X, Y, R', and R''. We find, for example, in the system the following series:—

Series of order $(n+2)$ $X' R' Y'$
 " " (n) $X R Y$
 " " $(n+2)$ $X'' R'' Y''$
 $R'' - R =$ nearly $R - R' =$ about 45.

Therefore, to find the properties of analogous compounds, we can establish proportions and find average values; the properties of all the elements are, properly speaking, in an intimate and mutual dependence. I call *atomanalogy* of an element the relation which joins R to X and Y, on the one hand, and to R' and R'' on the other hand. Thus As and Br on the one hand, and S and Te on the other, are the atomic analogues of Se, whose atomic weight is the medium one, viz.,—

$$78 = \left(\frac{75 + 80 + 32 + 125}{4} \right);$$

and, for example, the properties of SeH_2 answer to this atomic weight, because SH_2 occupies the middle place between AsH_3 — BrH and SH_2 — TeH_4 , &c. It is only in the series and in the ends of groups that the atomalogies are not quite valid, although we can there also see clearly the reciprocal relations, which can be shown conditionally by the arithmetical proportions, for example—

$$\mathbf{X}' : \mathbf{X} = \mathbf{R}' : \mathbf{R} = \mathbf{Y}' : \mathbf{Y} \text{ or } \mathbf{X}' : \mathbf{R}' = \mathbf{X} : \mathbf{R} = \mathbf{X}'' : \mathbf{R}''.$$

These reciprocal relations, to which the system founded on the periodic law leads us, furnish the possibility of explaining many isolated and doubted facts.

Different opinions have been given on the position of beryllium in the system since the researches of Awdceeff. He gives to the oxide the formula of magnesia; besides, glucina is very like magnesia in its properties. The periodic law gives the following proofs in favour of the formula BeO . If we adapted the formula Be_2O_3 , the atomic weight of beryllium would be $39.4 = 14.1$. In this case this element could not enter the system, because it would be by the side of nitrogen, and would have to possess very acid properties, and also to form higher oxides, having as formulæ Be_2O_5 and BeO_3 , of which it is not capable. If, on the contrary, we adopt BeO as the formula for its oxide, and 9.4 for the atomic weight of the metal, this element, conforming to the formula of oxides and to all its properties, would be between $L=7$ and $B=11$. The following proportions will suffice to show what I mean:—

1st. $\text{Be} : \text{Li} = \text{B} : \text{Be}.$

In effect the basic properties are more feebly shown in BeO than in Li_2O , and much more feebly in Be_2O_3 than in BeO . Chloride of beryllium is more volatile than chloride of lithium, and much more volatile than chloride of boron.

2ndly. $\text{Be} : \text{Mg} = \text{Li} : \text{Na} = \text{B} : \text{Al}.$

Oxide of beryllium is a less energetic base than MgO . Again, Li_2O reacts more feebly than Na_2O , and B_2O_3 more feebly than Al_2O_3 . This is why glucina dissolves in KHO .

The salts of BeO and of MgO do not present an absolute isomorphism; occasionally they show considerable differences in their crystalline forms. This fact is without importance, because the salts of Li and of Na and the compounds of B and of Al are none of them completely isomorphous.

Fluoride of beryllium is soluble in water, fluoride of magnesium is not; fluoride of boron is soluble, and fluoride of aluminium is not.

3rdly. $\text{Be} : \text{Al} = \text{Li} : \text{Mg} = \text{B} : \text{Si}$.

If, in spite of the difference of the formulæ of the oxides, oxide of beryllium corresponds in many respects to aluminium, Li_2O and MgO will also possess analogous properties; the same is true for B_2O_3 and SiO_2 . If, still further, the volume of an equivalent of BeO equal to—

$$\frac{25.4}{3.05} = 8.3,$$

approaches in volume an equivalent of aluminium equal to—

$$\frac{102.6}{4.0} = 8.5,$$

this concordance is repeated for the corresponding compounds of Li and $\frac{1}{2}\text{Mg}$, as well as for $\frac{1}{2}\text{B}_2\text{O}_3$ compared with $\frac{1}{2}\text{SiO}_2$.

The volume of $\text{LiCl} = 21$; therefore the volume of $\text{Li}_2\text{Cl}_2 = 42$, which corresponds to $\text{MgCl}_2 = 44$; the volume of $\text{BCl}_3 = 87$; therefore that of $\frac{1}{2}\text{BCl}_3 = 58$, which corresponds to $\frac{1}{2}\text{SiCl}_4 = 56$ ($\text{SiCl}_4 = 112$); the volume of $\text{B}_2\text{O}_3 = 39$; therefore the volume of $\frac{1}{2}\text{B}_2\text{O}_3 = 19.5$; whilst the volume of SiO_2 (amorphous) = 27, and, therefore, $\frac{1}{2}\text{SiO}_2 = 13.5$.

Again, does not Rose's observation, relative to the proximity which the volumes of equivalents of glucinum and of aluminium present, show the concordance which exists between the formulæ of the oxides. If BeO crystallises in the characteristic forms for Al_2O_3 , ZrO_2 crystallises in the same manner.

Therefore, the position of Be in the system would be elucidated on all points, so that we could form an opposition group to Li, Na, K, Rb, Cs, viz., the group Be, Mg, Ca, Sr, Ba, parallel in all respects.

In the same way the position of B relatively to C, Si, Al is indicated by what precedes. So I think it is sufficient to make the following proportions:— $\text{B} : \text{Al} = \text{Be} : \text{Mg}$; $\text{B} : \text{C} = \text{Be} : \text{B}$; $\text{B} : \text{Si} = \text{Be} : \text{Al}$. We can still add $\text{B} : \text{P} = \text{C} : \text{S}$, as it appears, if we examine the tables, and if we consider that B furnishes the compounds BCl_3 , B_2O_3 , BH_3O_3 , which correspond to PCl_3 , P_2O_3 , PO_3H_3 , &c., these circumstances coincide with those that we observe relative to CO_2 , C_2 , C_2H_2 , CH_2O_2 , and to SO_2 , S_2H_2 , SH_2O_2 .

(To be continued.)

DR. ANGUS SMITH'S REPORT

AS

INSPECTOR UNDER THE ALKALI ACTS.

RARELY have we met with a "Blue Book" at once so interesting and so satisfactory as Dr. Angus Smith's recent reports as Inspector under the Alkali Acts;—interesting from the importance of the subject dealt with, and satisfactory because the object of the Acts is being successfully carried out at a *minimum* of annoyance and trouble to all concerned. Our first reflection after examining the ensuing pages is, that Dr. Angus Smith still continues to approve himself "the right man in the right place." His thorough knowledge of industrial chemistry, his suggestive and fruitful mind, his tact, courtesy, and patience, his clear oversight of the whole field of operations, and of the rival interests to be dealt with, cannot be too highly

esteemed, and have won for him golden opinions among all competent judges. His method of leading instead of driving the chemical manufacturers, and of substituting, whenever possible, remonstrance and advice for summonses and fines, is perhaps slow, but it is sure. We know well that a part of the British public value remedial agencies and reforms of every kind simply in proportion to the disturbance they create,—a preference which, by the way, is the very sheet-anchor of quackery. Dr. R. A. Smith remarks that:—"Some of the public would have preferred to see the inspectors frequently in court with cases of complaint, but I know well that information must grow, and habits must grow, and that to torment men into doing what required much time to learn, was to return to the old system of teaching by the cane instead of through the intellect."

As to the success attained in working the Act, we learn that while the law allows 0.2 grain of muriatic acid to escape per cubic foot of mixed gases, from many works the escape has passed into the second decimal place, and in No. 2 and 3 districts this is the case for the yearly average. Hence the inspectors are now able to show—what formerly was a mere guess,—that there are much more sulphur acids passing into the atmosphere than muriatic acid. There are chimneys using coal alone, *i.e.*, where nothing but the products of the combustion of fuel escape, and where the acidity is yet above 0.2, and in Widnes it is sometimes above 1.0. Hence it is idle to complain that in all South Lancashire there is a lowering of the tone of vegetation. We have often sought to show that were all the chemical works of Widnes and St. Helens swept away, so long as the coal is so rich in sulphur, beautiful trees are out of the question. We may further ask whether the prosperity of our manufactures is not, after all, of at least as much importance as the growth of trees in districts that never possessed any attractive scenery?

The attention of the inspectors has not been by any means confined to the escape of muriatic acid, sulphur, and nitrogen acids from chimneys and chambers. They have also expended much research on the gases evolved from the waste-heaps. Two objects have here to be kept in view; the abatement of the nuisance and the utilisation of the sulphur, now chiefly wasted. The escape of sulphuretted hydrogen, Dr. Angus Smith thinks, will never be entirely done away with till the Le-Blanc process is laid aside. This, again, is a consummation scarcely to be expected until "we learn to make ammonia from the nitrogen of the air and the hydrogen of water." In the meantime, a variety of methods have been proposed to minimise the evil. Dr. Mond seeks to oxidise the sulphur to a certain extent by blowing air through the mass fresh from the tanks and when the exact amount of hyposulphurous acid is formed capable of destroying the sulphide of hydrogen or of calcium remaining, he dissolves the soluble matter in water, and adds hydrochloric acid, which releases both the acids, so as to let them act on each other, when free sulphur is thrown down. This process has been used by Mr. Worsley, of Netham, near Bristol, who finds it remunerative.

At the Alkali Works of Dieuze any salts of iron are mixed with the mass, which is frequently stirred, so that the iron is alternately sulphurised and oxidised. At each sulphuration hyposulphite and bisulphide are formed. The two are mixed in proper proportions, so that on the addition of hydrochloric acid all the sulphur may be thrown down.

Dr. Smith lays before the public the outline of a new process which seems very promising, and which we should like to see tried on the large scale. If MnO_2 is added to liquid waste the sulphur is thrown down rapidly, and the calcium is oxidised. On exposing this manganese sulphide to air it is oxidised and sulphide is thrown out in a free state. If air is passed through the liquid in which the sulphide of manganese lies oxidation takes place and oxide of manganese is formed; part of the sulphur is at the same time converted into hyposulphurous acid. The

use of hot water at high pressures, as proposed by Krause, is the most scientific method of dealing with solid waste, if not too costly.

Sulphuretted hydrogen is now successfully burnt to sulphuric acid on the large scale by means of a process devised by the late Mr. W. Hunt, of Wednesbury, and now carried out by Mr. P. Spence, of Manchester, at the Frizinghall Works, near Bradford.

The manufacture of coke, as at present conducted, involves waste and nuisance on a gigantic scale. We learn that the ammonia thus wasted in 1876 equalled 139,764 tons of sulphate of ammonia, representing a value of £2,795,280, without taking into account the coal-tar likewise sacrificed. Dr. Angus Smith is of opinion that every effort should be made to prevent this destruction, and recommends the coke-ovens of MM. Carvès and Co. to all who use coals containing from 25 to 33 per cent of volatile matter. The notion that it is impossible to produce a dense coke and to utilise the by-products is merely an apology for carelessness.

The author's remarks on the importance of securing the ammonia now wasted should in these times commend themselves to a much wider circle than merely to chemical manufacturers. It is to be noted that a great coal-owner asked Dr. Angus Smith, about thirty years ago, to make experiments on the best mode of saving the ammonia from the coke-furnaces, but was not willing to pay a shilling towards the expense of operations which must necessarily be conducted on a large scale. Upon which, says Dr. Smith, "I gave up the idea." Such is the general way in which chemists are treated.

We cannot too strongly commend this report to the notice of manufacturing chemists. In addition to other valuable information they will find in it the substance of several inventions which the author might legitimately have patented, but which he freely publishes for the benefit of the trade.

THE TESTING OF PETROLEUM ACT, 1879.

The Home Office has just issued the following instructions as to the mode of testing petroleum so as to ascertain the temperature at which it will give off inflammable vapour:—

The oil cup consists of a cylindrical vessel 2 inches diameter, 2½ inches height (internal), with outward projecting rim 5-10th inch wide, ½ inch from the top, and 1½ inch from the bottom of the cup. It is made of gun-metal or brass (17 B.W.G.), tinned inside. A bracket, consisting of a short stout piece of wire bent upwards and terminating in a point, is fixed to the inside of the cup to serve as a gauge. The distance of the point from the bottom of the cup is 1½ inches. The cup is provided with a close-fitting overlapping cover made of brass (22 B.W.G.), which carries the thermometer and test-lamp. The latter is suspended from two supports from the side by means of trunnions, upon which it may be made to oscillate; it is provided with a spout, the mouth of which is 1-16th inch in diameter. The socket which is to hold the thermometer is fixed at such an angle and its length is so adjusted that the bulb of the thermometer when inserted to its full depth shall be 1½ inches below the centre of the lid. The cover is provided with three square holes, one in the centre, 5-10th inch by 4-10th inch, and two smaller ones, 3-10th inch by 2-10th inch, close to the sides and opposite each other. These three holes may be closed and uncovered by means of a slide moving in grooves, and having perforations corresponding to those on the lid. In moving the slide so as to uncover the holes, the oscillating lamp is caught by a pin fixed in the slide, and tilted in such a way as to bring the end of the spout just below the surface of the lid. Upon the slide being pushed back so as to cover the holes the lamp returns to its original position. Upon the cover in front

of, and in line with, the mouth of the lamp is fixed a white bead, the dimensions of which represent the size of the test-flame to be used. The bath or heated vessel consists of two flat-bottomed copper cylinders (24 B.W.G.), an inner one of 3 inches diameter and 2½ inches height, and an outer one of 5½ inches diameter, and 5½ inches height; they are soldered to a circular copper plate (20 B.W.G.) perforated in the centre, which forms the top of the bath, in such a manner as to enclose the space between the two cylinders, but leaving access to the inner cylinder. The top of the bath projects both outwards and inwards about ½ inch; that is, its diameter is about ½ inch greater than that of the body of the bath, while the diameter of the circular opening in the centre is about the same amount less than that of the inner copper cylinder. To the inner projection of the top is fastened, by six small screws, a flat ring of ebonite, the screws being sunk below the surface of the ebonite to avoid metallic contact between the bath and the oil-cup. The exact distance between the sides and bottom of the bath and of the oil-lamp is ¼ inch. [This statement relates to the distance between the sides and bottom of the cup and the walls of the inner cylinder which forms the air-chamber.] A split socket similar to that on the cover of the oil-cup, but set at a right angle, allows a thermometer to be inserted into the space between the two cylinders. The bath is further provided with a funnel, an overflow pipe, and two loop handles. The bath rests upon a cast-iron tripod-stand, to the ring of which is attached a copper cylinder or jacket (24 B.W.G.) flanged at the top, and of such dimensions that the bath, while firmly resting on the iron ring, just touches with its projecting top the inward-turned flange. The diameter of this outer jacket is 6½ inches. One of the three legs of the stand serves as support for the spirit-lamp attached to it by means of a small swing bracket. The distance of the wick-holder from the bottom of the bath is 1 inch. The lamp is filled through the funnel. In both thermometers the capillary tube is widened at the top to prevent breakage through over-heating. The line on the scale of the long-bulb thermometer indicating 130° is rendered conspicuous by being drawn across the whole width of the ivory back. In a similar manner the line indicating 73° is specially marked on the round bulb thermometer. Two thermometers are provided with the apparatus, the one for ascertaining the temperature of the bath, the other for determining the flashing-point. The thermometer for ascertaining the temperature of the water has a long bulb and a space at the top. Its range is from about 90° to 190° F. The scale (in degrees of Fahrenheit) is marked on an ivory back fastened to the tube in the usual way. It is fitted with a metal collar, fitting the socket, and the part of the tube below the scale should have a length of about 3½ ins., measured from the lower end of the scale to the end of the bulb. The thermometer for ascertaining the temperature of the oil is fitted with collar and ivory scale in a similar manner to the one described. It has a round bulb, a space at the top, and ranges from about 55° to 150° F.; it measures from end of ivory back to bulb 2½ inches.

A model apparatus is deposited at the Weights and Measures Department of the Board of Trade.

DIRECTIONS FOR APPLYING THE FLASHING TEST.

1. The test apparatus is to be placed for use in a position where it is not exposed to currents of air or draughts.
2. The heating-vessel or water-bath is filled by pouring water into the funnel until it begins to flow out of the spout of the vessel. The temperature of the water at the commencement of the test is to be 130° F., and this is to be attained in the first instance either by mixing hot and cold water in the bath, or in a vessel from which the bath is filled, until the thermometer which is provided for testing the temperature of the water gives the proper indication; or by heating the water with the spirit-lamp (which is attached to the stand of the apparatus) until the required temperature is indicated. If the water has been heated too highly, it is easily reduced to 130° by pouring

in cold water little by little (to replace a portion of the warm water) until the thermometer gives the proper reading. When a test has been completed, this water-bath is again raised to 130° by placing the lamp underneath, and the result is readily obtained while the petroleum-cup is being emptied, cooled, and re-filled with a fresh sample to be tested. The lamp is then turned on its swivel from under the apparatus, and the next test is proceeded with.

3. The test-lamp is prepared for use by fitting it with a piece of flat plaited candle-wick, and filling it with colza or rape oil up to the lower edge of the opening of the spout or wick-tube. The lamp is trimmed so that when lighted it gives a flame of about 0.5 of an inch diameter, and this size of flame, which is represented by the projecting white bead on the cover of the oil-cup, is readily maintained by simple manipulation from time to time with a small wire trimmer. When gas is available it may be conveniently used in place of the little oil-lamp, and for this purpose a test-flame arrangement for use with gas may be substituted for the lamp.

4. The bath having been raised to the proper temperature, the oil to be tested is introduced into the petroleum-cup, being poured in slowly. In pouring in the oil to be tested great care should be taken not to splash it against the sides of the cup until the level of the liquid just reaches the point of the gauge which is fixed in the cup. In warm weather the temperature of the room in which the samples to be tested have been kept should be observed in the first instance, and if it exceeds 65° the samples to be tested should be cooled down (to about 60°) by immersing the bottles containing them in cold water, or by any other convenient method. The lid of the cup, with the slide closed, is then put on, and the cup is placed into the bath or heating-vessel. The thermometer in the lid of the cup has been adjusted so as to have its bulb just immersed in the liquid, and its position is not under any circumstances to be altered. When the cup has been placed in the proper position the scale of the thermometer faces the operator.

5. The test-lamp is then placed in position upon the lid of the cup, the lead-line or pendulum, which has been fixed in a convenient position in front of the operator, is set in motion, and the rise of the thermometer in the petroleum-cup is watched. When the temperature has reached about 66° the operation of testing is to be commenced, the test-flame being applied once for every rise of one degree in the following manner:—The slide is slowly drawn open while the pendulum performs three oscillations, and is closed during the fourth oscillation.

If it is desired to employ the test apparatus to determine the flashing-points of oils of very low volatility, the mode of proceeding is to be modified as follows:—The air-chamber which surrounds the cup is filled with cold water to the depth of 1½ inches, and the heating-vessel or water-bath is filled as usual, but also with cold water. The lamp is then placed under the apparatus, and kept there during the entire operation. If a very heavy oil is being dealt with the operation may be commenced with water previously heated to 120° instead of with cold water.—*Chemist and Druggist.*

CRYSTALLISATION OF CARBON.

At a meeting of the Glasgow Philosophical Society on Wednesday, Mr. James Maclear, of the St. Rollox Chemical Works, stated that on the 12th of this month he had sent a communication to Mr. Dixon, the secretary, giving the ascertained results of some discoveries he had made, but they were of such a startling nature that he thought it proper to have the note kept sealed until he had satisfied himself by all means in his power, and obtained more competent opinions than his own, as to the nature of these results. Since that time he had been in London and had shown the results to Profs. Tyndall, Warrington Smyth, and others, and they were now in the hands of Mr. Maskelyne, of the British Museum. They were briefly these:

—After having thought carefully from time to time over the subject, and made many abortive experiments extending over a period which dated back to 1866, he had at last succeeded in obtaining crystalline forms of carbon. They were perfectly pure and transparent, and had all the refractive power of diamonds. They had the crystalline form of diamonds, and resisted acids, alkalies, and the intense heat of the blowpipe. They also scratched glass, and the only other test that remained was as to whether they would scratch diamonds or be scratched by them, and as to the refractive index of the crystal itself and the measurement of the angle of the crystals. These tests, as he had said, had not been carried out, but they would be shortly, and he hoped to put some of the specimens before the society on a future occasion. He had no doubt in his own mind, and neither was there any doubt in the minds of the scientific gentlemen whom he had consulted, that they were diamonds, but in the meantime he preferred to describe them as pure crystalline forms of carbon. The measurement of the angle of the forms he had obtained was 1.32nd part of an inch.

Just as we were going to press we received a telegram from Mr. Maclear, in which he says that he has been working on the subject about thirteen years with varying success. He did not announce the discovery, however, until absolutely certain, and reserves details of the processes for the present. There is no difficulty in producing carbon crystals, apparently diamond, from any carbon compound whatever and by a variety of processes based on a great physical fact. Even combustion products are quite suitable. Mr. Maclear has promised us specimens for physical and chemical examination.

ON BESSEMER STEEL PLATES.

By SERGIUS KERN, M.E., St. Petersburg.

THE author some time ago (CHEMICAL NEWS, vol. xxxviii, No. 966,) remarked that for the rolling of boiler plates out of Bessemer ingots, it is preferable to use ingots, hammered after casting. This method is employed in many works, and the reasons for doing so may be explained as follows:—

1. The plates obtained by the rolling of hammered ingots have a smooth fine surface. Flaws, scale, or excavations are seldom observed, and if they happen, this must be attributed to imperfect rolling; because an ingot which hammers well, and shows no defects during this process, is likely to roll well into plates.

2. Plates out of hammered ingots have a higher density, good structure, and are more uniform in their mechanical qualities. Such plates, even unannealed, will always stand the test in the limits of the Lloyd's regulations.

On the other hand, plates rolled directly out of unhammered ingots, show a great fluctuation in their mechanical qualities. This may be seen by the examination of the annexed tables. A want of uniformity in the results of the Table A will be observed.

AVERAGE ANALYSES.

1. Plates Rolled out of Hammered Ingots.

	Per cent.
Carbon	0.20
Silicon	0.02
Phosphorus	0.04
Manganese	0.32
Sulphur	0.03
Copper	none

2. Plates Rolled out of Unhammered Ingots.

	Per cent.
Carbon	0.24
Silicon	0.02
Phosphorus	0.03
Manganese	0.30
Sulphur	0.03
Copper	none

TABLE A.

Unannealed Plates, prepared out of Unhammered Ingots.

No. of Samples.	Tensile Strength Per Square Inch. (In Tons.)	Elongation. (Per Cent.)
1.	25.75	31.00
2.	31.24	22.50
3.	29.17	13.25
4.	32.28	17.50
5.	29.27	25.25
6.	32.50	21.50
7.	31.48	23.50
8.	32.23	22.00
9.	32.64	25.00
10.	30.18	17.87
11.	32.50	19.50
12.	27.94	25.37

Size of specimens: length 8 inches, breadth 1.5 inches, thickness 0.625 inch.

TABLE B.

Unannealed Plates, prepared out of Hammered Ingots.

No. of Samples.	Tensile Strength per square inch. (In tons.)	Elongation. (Per cent.)
1.	27.82	25.50
2.	28.00	22.50
3.	27.00	22.00
4.	27.70	28.50
5.	27.56	26.25
6.	27.48	30.00
7.	27.12	25.12
8.	27.55	25.50
9.	28.00	23.50
10.	27.78	26.75
11.	27.12	25.00
12.	27.55	26.25

of specimens: as mentioned in Table A.

The hammered and unhammered ingots were rolled into plates under the same conditions, and were heated to a white non-welding heat. In most cases the ingots were finished into plates from one heat.

PROCEEDINGS OF SOCIETIES.

CHEMICAL SOCIETY.

Thursday, December 18, 1879.

Mr. WARREN DE LA RUE, President, in the Chair.

A BALLOT for the election of Fellows was held, Dr. Messell and Mr. Neison being appointed scrutators. The following gentlemen were declared duly elected:—G. S. Allbright, J. R. Ashwell, T. Blackhouse, E. Buckney, J. Bemrose, W. J. F. Churchouse, M. Cochrane, E. J. Day, W. K. Dunstan, E. Francis, F. Hatton, J. Howard, J. J. Hummell, R. E. Holloway, W. R. Eaton Hudgkinson, T. S. Humpidge, E. Hughes, R. Jones, J. Knowles, A. Leibius, H. F. Morley, E. F. Mondy, H. Newton, J. Parette, J. W. Smith, J. Steiner, J. Snodgrass, A. Scott, G. Stallard, J. M. Wilson. The following certificates were read for the first time:—W. Macnab, W. B. Roberts, G. Salet, T. Terrell.

The SECRETARY then read a paper, "On the Specific Volume of Crystallisation," by T. E. THORPE and J. J. WATTS. Some years ago Playfair and Joule pointed out that the volumes of certain highly hydrated salts, as for example, sodium carbonate with 10 molecules of water, also the alkaline arsenates and phosphates with 12 mole-

cules, are equal to that of the water, considered as ice, which they respectively contain: thus the molecules of the salt seem to exist in the interstitial spaces of the ice. This law does not hold good for salts less highly hydrated. Thus, in borax, sodium pyrophosphate, and the normal aluminium sulphate the volume seems to be made up of the water considered as ice, together with that of the base as existing in the free state. Schiff has shown that members of certain classes of hydrated salts have the same specific volume. Thus all the alums have a specific volume 277, double sulphates, $M_2M''(SO_4)_2$, 207, and the vitriols, $M''SO_4 \cdot 7H_2O$, 146. The authors have determined the precise relation between the specific volumes of various sulphates of copper, magnesium, zinc, nickel, cobalt, iron, and manganese, and their respective degrees of hydration. They have incorporated in the present paper some results placed at their disposal by Dr. Playfair. Details of the preparation and analysis of the various hydrates employed are given. The specific gravity was determined by weighing in benzene at 15° C. The results are contained in the following table:—

Hydrate.	0.	1.	2.	3.	4.	5.	6.	7.
CuSO ₄ ..	44.4	54.3	67.0	80.0	—	109.1	—	—
MgSO ₄ ..	44.8	55.6	67.0	—	—	112.4	130.8	146.8
ZnSO ₄ ..	45.6	54.7	66.6	—	—	113.7	130.2	146.8
NiSO ₄ ..	44.6	56.5	—	—	—	—	129.0	144.6
CoSO ₄ ..	44.7	55.2	70.9	—	97.4	114.6	130.1	146.0
MnSO ₄ ..	45.0	55.7	73.6	86.6	98.2	114.4	—	—
FeSO ₄ ..	44.5	56.2	67.7	—	100.5	—	—	146.7
Means ..	44.8	55.5	68.8	83.3	93.7	112.9	130.0	146.2

From these results it appears that in the case, at least, of the so-called magnesian sulphates the volumes occupied by the several molecules of water varies with the degree of hydration. The first molecule, the "constitutional water" of Graham, occupies less bulk than any other. Its mean relative value is 10.7. The value of the second is 13.3; of the third, 14.5; of the fourth, 15.4; of the fifth (taking the mean of the most concordant numbers, ZnSO₄, CoSO₄ and MnSO₄), 15.6; of the sixth, 15.7; of the seventh, 16.2. These results accord with the fact that the different molecules of water in a hydrated salt are held with various degrees of tenacity, as shown by the different intensities of heat needed to expel them, the amount of energy required standing in some relation to the degree of condensation in the combined molecule. The authors point out the importance of estimating the amounts of heat resulting from the combination of successive molecules of water with the different sulphates. Graham has already shown that more heat was evolved in the combination of the first molecule than in that of any of the remaining molecules. In other words, that the amount of heat developed is related to the degree of condensation of the combined molecules. As the foregoing numbers express the volumes in cubic centimetres of the equivalent of the salts in grammes, it appears that equivalent quantities of these different sulphates occupy respectively the same volume in space, or, in other words, the unit volume contains the same number of molecules of the different salts.

Prof. McLEOD then read a "Note on the formation of Ozone during the Slow Oxidation of Phosphorus." The active substance formed during the slow oxidation of phosphorus is probably either ozone or peroxide of hydrogen. The latter substance is readily destroyed by alkalies, a solution of chromic acid, or a solution of alkaline permanganate, whilst ozone is unaffected either by a solution of sodic carbonate or by chromic acid, and appears to be only slightly attacked by alkaline permanganate. Air in which phosphorus was slowly oxidising was drawn through a U-tube 9½ inches long (filled with fragments of glass, containing in succession sodic carbonate saturated with carbonic anhydride, a mixture of potassic dichromate and sulphuric acid, and potassic permanganate previously

saturated with carbonic anhydride), and then into a flask containing a solution of potassic iodide and starch. In all cases the latter solution became blue, both when the U-tube was cold and when heated to 100° . Similar results were obtained when a U-tube $12\frac{1}{2}$ inches long was used, packed with small pieces of pumice and saturated with solution of sodic carbonate. The effect of heat on the gas was tried. The gas was aspirated through a narrow U-tube, which was heated to 150° and 200° . Beyond this U-tube were placed, first, a weighed U-tube packed with pumice and sulphuric acid, and, secondly, a flask with solution of potassic iodide and starch acidified with sulphuric acid. The U-tube was weighed before and after each experiment, and the blue solution titrated with decinormal sodic thiosulphate. The gas was aspirated at the rate of 1 litre per hour. The following results were obtained:—

Gas Aspirated.	Temp. of U-tube.	Increase of Sulphuric Acid Tube.	Thiosulphate Used.
4600 c.c.	cold	0.0026 grm.	2.55 c.c.
2760 "	100°	0.0008 "	1.9 "
4600 "	150°	0.0026 "	3.2 "
2760 "	200°	0.0006 "	1.8 "

1 c.c. of thiosulphate = 0.017 grm. of hydroxyl, which on decomposition forms 0.009 grm. of water; and as at least one half of the hydroxyl might be assumed to be decomposed, an increase of the sulphuric acid tube in the last experiment should be 0.016 grm. instead of only 0.0006. Hydroxyl combines with acids. The gas from phosphorus was exposed to the action of strong sulphuric acid for four days without losing its activity. It is extremely improbable that ozone and hydroxyl are both formed, as these substances destroy each other. The author therefore concludes that the gas obtained during the slow oxidation of phosphorus possesses the properties of ozone, and not those of hydroxyl, the only known peroxide of hydrogen.

Mr. KINGZETT had listened to the paper with great interest. Up to the present time no evidence had been put on record to prove that this gas from phosphorus contained ozone, and as he had shown that certain hydrocarbons which had been supposed to produce ozone really formed hydroxyl, he had assumed, in the absence of any evidence, that hydroxyl was formed in this case. After referring to the researches of Cornu, Mr. Kingzett said that he could confirm the observation of Prof. McLeod that peroxide of hydrogen and ozone in the presence of an acid did not decompose each other.

After some remarks from the President, Prof. McLeod briefly replied.

Mr. W. H. PERKIN then read a paper "*On the Analysis of Organic Bodies containing Nitrogen.*" Some years since the author wished to determine the halogens as well as the carbon and hydrogen in bodies containing nitrogen. The substance was burnt in oxygen, and the products of combustion passed over a weighed quantity of pure metallic silver. The difficulty arose as to how to get rid of the nitrous fumes: plumbic peroxide was first tried between the water absorbing apparatus and potash bulbs. This succeeded as far as the carbon was concerned, but as nitrous fumes were absorbed by the sulphuric acid the hydrogen determinations were too high. Plumbic peroxide was then placed in the combustion-tube, but great difficulty was experienced in heating it to just the proper temperature. Potassic chromate was then tried, and was found to succeed admirably: it absorbed the nitrous fumes completely, had no action on carbonic anhydride, and could be heated without fear. The author therefore recommends the use of about 6 to 7 inches of potassic chromate in the combustion of all substances containing nitrogen instead of freshly reduced copper, which is very hygroscopic and occludes hydrogen, or of silver, which requires an inconveniently high temperature. The potassic chromate should be free from an excess of alkali: a trace of bichromate is not harmful, though bichromate does not work

so well as chromate. The chromate should be roughly powdered or granulated by evaporating its solution to dryness with constant stirring; or, still better, coarsely powdered pumice saturated with solution of chromate may be used. The chromate should be kept at a scarcely dull red-heat. The author does not state how many times the chromate can be used. Chromate also absorbs sulphurous acid. It does not retain iodine vapour, but would be probably useful with substances containing chlorine or bromine.

After a few remarks from Mr. Warrington the Society adjourned to January 15.

PHYSICAL SOCIETY.

Ordinary Meeting, December 13, 1879.

Prof. W. G. ADAMS in the Chair.

NEW MEMBERS:—Mr. J. H. Poynting, Mr. R. T. Glazewood, Dr. R. C. Shettle, Prof. Rowland, Mr. John Gray, D.Sc., Mr. H. R. Brook, Mr. E. B. Sargent, Mr. E. Paterson.

"On the Graduation of the Sonometer," by Mr. J. H. POYNTING, Trinity College. The author had endeavoured to reduce the present arbitrary readings of the Sonometer of Prof. Hughes to absolute measure by adapting the formula given in Maxwell's "Electricity" (vol. ii., chap. xiv.), to the induction of two circular coils on the same axis, but separated by a distance greater than the radii of the coil, on a third coil intermediate. By applying the formula thus obtained to the results of Prof. Hughes for different metals, he finds that either the specific resistances of metals as given in the tables are not the same as the resistances of the metals employed by Prof. Hughes, or that the induction effect of the balance or sonometer is not proportional to the conductivity of the metal.

Prof. AYRTON reminded the Society that at a former meeting he had shown mathematically that the effect was not proportional to the conductivity, but to an exponential function of the conductivity.

Mr. CHANDLER ROBERTS, F.R.S., stated that Prof. Hughes did not profess that the metals used by him to obtain his results were pure.

Prof. ADAMS mentioned that Prof. Hughes had shown that the effect was dependent on other conditions than the mere purity of the metal.

Dr. J. A. FLEMING, St. John's College, Cambridge, exhibited and described a new form of Wheatstone Balance, designed principally for comparing the B.A. units of resistance deposited in the Cavendish Laboratory. The divided resistance is a circular platinum-iridium wire, and an arm fitted with a contact at its extremity revolves round after the manner of circular resistance-coils, thus altering the ratio of the divided resistances. The contact is a knife-edge of platinum, and it is "made" and "broken" by hand as with a key. A series of ingenious copper-mercury cups are fitted to the balance so as to permit of two coils being compared at any temperature with great exactness by the method suggested by Prof. Foster and adopted by Prof. Crystal, of Cambridge. This consists in exchanging the positions of the units on the balance, and observing the difference in the results. By Dr. Fleming's arrangement this exchange can be effected without removing the coils from heating apparatus in which they are placed, or otherwise altering their conditions. The mercury contact-cups and the heating-cans were also improved by Dr. Fleming for the purpose of facilitating accuracy of results.

Prof. PERRY described a Dispersion Photometer devised by himself and Prof. Ayrton for the purpose of comparing intense lights, such as the electric, with a standard candle without necessitating much room in order to put the stronger light at a distance from the screen proper to give an illumination equal to that of the candle. To reduce the distance of the stronger light from the screen, the authors had inserted a lens in the track of the beam, so as to disperse the beam to a degree which could be determined by an easy formula. Thus, by artificially diluting the power

ful beam they could compare it with the feebler beam from the standard light in a shorter space. For an electric light of 6400 candles only 8 feet need be required by the new plan instead of 80 feet by the unassisted method.

Dr. JOHN HOPKINSON, F.R.S., stated that he had actually used the same method for some months past in his electric light experiments, and recommended a plano-convex lens as the best to use, and suggested that the focal length should be calculated. He thought that the error due to absorption could easily be obviated.

Prof. AYRTON then described a method by which Prof. Perry and he had determined the value of g , or the coefficient of gravity, at the Imperial Engineering College, Tokio, Japan, by means of pendulums. Their result is 980.06; and calculation from the position of the place makes it 979.8.

An Improved Form of Spherometer, designed by Mr. W. Goolden, and made by Mr. Adam Hilgar was exhibited to the meeting. The frame is of aluminium, combining lightness and rigidity; the legs and screws of hard steel. The screw carries a drum divided into 1000 parts, and the instrument gives a reading to the $\frac{1}{1000}$ of an inch by the usual method of touch. Increased sensitiveness is got by employing a galvanometer to indicate the contact of the middle pointer with the surface. By this means it is made correct to $\frac{1}{100000}$ inch. A smaller form of the instrument, correct to $\frac{1}{10000}$ inch, was also shown.

MANCHESTER LITERARY AND PHILOSOPHICAL SOCIETY.

Ordinary Meeting, November 18, 1879.

J. P. JOULE, D.C.L., LL.D., F.R.S., &c., President, in the Chair.

THE PRESIDENT noticed the lamented death, since the last meeting, of Prof. James Clerk-Maxwell, an event which, having occurred in the prime of his life and in the midst of his usefulness and his splendid researches, was felt most severely by the Society and the whole scientific world.

"Recording Sunshine," by DAVID WINSTANLEY, F.R.A.S. So far as I have seen there is in use at present but one form of apparatus which effects an automatic registration of the duration and the times of sunshine, and that is the instrument of Campbell, in which a sphere of glass is so disposed as to burn a piece of wood or paper by the concentration of his rays when the sun may chance to shine. During the past few years I have devoted some attention to this matter and devised a number of appliances having the same object for their end but differing materially both in their construction and in the manner of their use from the apparatus I have named.

One of these, with your permission, I will now describe.

It is an arrangement which places a lead pencil on a sheet of paper and writes down therewith when and for how long the sunshine lasts.

It consists essentially of a differential thermometer with a long horizontal stem, in which latter is contained throughout the greater portion of its length some fluid intended to operate by its weight. This thermometer is attached to a scale beam or some equivalent device which also carries the pencil by means of which the record shall be made.

The whole is so arranged that in its normal state it rests gently—upon that side to which the pencil is *not* attached—on an embankment provided for that end.

Close beneath the pencil point a disc of metal rotated at the proper speed carries a paper dial whereon marks and figures are engraved corresponding with the hours at which the sun may shine.

When using this instrument I have it enclosed within a box which permits one bulb only of the thermometer—that most distant from the clock—to be affected by the

radiance of the sun, which when it hines expands the air contained therein, forces the fluid along the tube, and by altering the equilibrium of the beam brings some portion of its weight to bear upon the pencil point, and so the record is commenced.

When the sun becomes obscured, the air expanded by his rays contracts, the fluid in the tube returns, the normal equilibrium is restored, and the pencil ceases to produce its mark.

In the instance of the instrument I use the stem of the thermometer is 18 inches long and the eighth of an inch or thereabouts in bore.

Mercury in consideration of its weight is the fluid I employ, and in conjunction with it some sulphuric acid is enclosed, because of the mobility which is thereby gained. I am aware that in these circumstances mercuric sulphate is very slowly formed, but after two years' lapse of time no inconvenience has been caused thereby and the mobility of the mercury remains.

The bulbs of the thermometer are 2 inches in diameter or thereabouts, and that they may be more rapidly affected the glass thereof is thin. Both are blacked, and the one intended to receive the radiance of the sun projects above the box in which the apparatus is contained into a dome of glass.

"On some Notices in Classical Authors of the Action of Sunlight on Purple Dye," by JAMES BOTTOMLEY, D.Sc., F.C.S.

"On the Origin of the Word Chemistry," by CARL SCHORLEMMER, F.R.S. Chemistry as a science is first mentioned* by Julius Maternus Firmicus, a native of Sicily, and procurator under Constantine the Great. He wrote at about 336 a work on Astrology, which has been preserved only in a defective state, and is commonly known by the name of *Mathesis*.

In this work he states that by observing the position of the moon, in respect to certain heavenly bodies or constellations, at the hour when a child is born, its future inclinations can be predicted. He continues: *Et si fuerit haec domus Mercurii, Astronomiam. Si Veneris, cantilenas et laetitiam. Si Martis, opus armorum et instrumentorum. Si Jovis, divinum cultum et scientiam in lege. Si Saturni, scientiam alchimiae. Si Solis, providentiam in quadri-pedibus. Si in Cancero, domus sua, scientiam dabit omnium quae exiunt de aqua.*†

Other editions of this work have also *scientia alchimiae*,‡ but Vossius informs us that in the manuscripts it is *chimiae*.§ He says: *Alchimia scientiam nominat Firmicus, lib. III., cap. XV. Ita quidem editum ab Aldo, sed in chirographis est chimia.*

Athanasius Kircher also states that the manuscript in the library of the Vatican has *chymia* and not *Alchymia*.||

Firmicus does not give any explanation of this term. However, another writer, who probably lived at the same time, if not earlier, explains it. Zosimus, the Panopolite, according to Georgios Synkellos, a writer of the ninth century, states that *χημεία* (or *χυμεία*, as some manuscripts have) meant the art of making gold or silver.¶

The curious passage in which this word occurs is the following:—

"The sacred Scriptures inform us that there exists a tribe of genii, who make use of women. Hermes mentions this circumstance in his Physics; and almost every writing (*λογος*), whether sacred (*φανερος*) or apocryphal, states the same thing. The ancient and divine Scriptures inform us that the angels, captivated by women, taught them all the operations of nature. Offence being taken at this, they remained out of heaven, because they had taught mankind all manner of evil, and things which could

* Kopp, "Beiträge zur Geschichte der Chemie," 43.

† Julius Firmicus de nativitatibus; Ed. Simon Bivillaqua, Venice, 1497.

‡ Ed. Aldus Manutius, Venice, 1499; Ed. Nicolaus Brucknerus, Bâle, 1533.

§ Etymologicon linguae latinae, Amsterdam, 1693.

¶ Kopp, loc. cit., 9.

|| Thomson's "History of Chemistry," 3.

not be advantageous to their souls. The Scriptures inform us that the giants sprang from their embraces. *Chema* is the first of their traditions respecting these arts. The book itself they called *Chema*; hence the art is called *Chemia*."

It is not difficult to trace the origin of this myth. We find it first in Genesis, chap. vi.: "And it came to pass, when men began to multiply on the face of the earth, and daughters were born unto them, that the sons of God saw the daughters of men that they were fair, and they took them wives of all which they chose."

"There were giants in the earth in those days; and also after that, when the sons of God came in unto the daughters of men, and they bare children to them, the same became mighty men, which were of old, men of renown."

Alluding to this later writers state that the fallen angels taught women all the secrets of nature.* That one of these is the art of making gold and silver is, however, first mentioned by Zosimus. Other Greek writers use the word *Chemia* or *Chymia* in the same sense; in print we find it first in the Lexicon of Suidas, who lived in the eleventh century, and defines *χημεία* as "the preparation of gold and silver."

All the earlier Greek writers who mention this word were in close connection with the university of Alexandria; from this it has been inferred that the artificial preparation of the noble metals was first attempted in Egypt.

That country was conquered by the Arabians in 640. Here they made undoubtedly their first acquaintance with chemical science; they prefixed their article to the Greek name, and thus introduced the terms, Alchemy, Alchimy, or Alchymy.

The origin and meaning of these terms have often been discussed. Plutarch states that the old name of Egypt was *χημεία*; that it was so called on account of its black soil, and that the same word designated the black of the eye. From this the conclusion has been drawn that chemistry originally meant the science of Egypt, or the black of the eye being the symbol of darkness and mystery, chemistry was the secret or black art. But alchemy has never been called the black art, a name which was exclusively reserved for magic or necromancy.

It has also been stated that the name was derived from the Arabic *kema*, to hide, while others have maintained that the founder of our science was Cham or Ham, the son of Noah, or an Egyptian king with the name of Chemmis. It has further been suggested that the name of the science was derived from *χέω*, to melt; or from *χυμός*, juice or liquid.

To this it has been objected that the original spelling was *χημεία* and not *χυμεία*, which, although Hermann Kopp, the great historian of chemistry, inclines to this view, has not yet been proved satisfactorily. Humboldt believes that the latter word got into some manuscripts by a mistake of the transcriber, and continues: "Alchemy commenced with the metals and their oxides, and not with the juices of plants." This objection, however, cannot be maintained at all, because vegetable juices, or, at least, substances designated by their names, are mentioned by the older alchemists as the most potent substance by which transmutations could be effected.†

Some time ago my friend Prof. Theodores called my attention to an interesting paper on this subject, published by Prof. Gildemeister,‡ in which he maintains the derivation of the word chemistry from *χυμός*. According to him *kimiyā* in Arabic does not originally have an abstract meaning, and is the name, not of a science, but of a body by which, or rather by a substance obtained from which, the transmutation of metals is effected; it is synonymous with *iksir*. Alchemy, as a science, was called: The preparation of *kimiyā* or *iksir*, also the science of the preparation of *kimiyā*, or, more shortly, science of *kimiyā*. In the Arabic Lexicon (Qāmūs) *al-iksir* is explained by *al-kimiyā*,

and the latter again by the former, or by any medium which, applied to a metal, transports it into the sphere of the sun or the moon, i.e., converts it into gold or silver.

Even to this day the word is used in the concrete sense; Kotschy* relates that the pasha of Nicosia talked much of flowers, chiefly *kimia*, a plant having the property of converting metals into gold.

The later writers, however, called the science shortly *al-kimiyā*, and retained the term *al-idsir* (elixir) for the transmuting medium or the philosopher's stone. This latter word is identical with *ἐξήριον*, as the writers of the Alexandrine school called the philosopher's stone,† while the same name was employed by the physicians for a healing powder, used for sprinkling over wounds, i.e., a desiccative powder (from *ξηρός*, dry).‡

Now the correlate to dry is moist or liquid, *χυμός*, and from this is derived *χυμεία*, a moist substance corresponding to *λίθια*, a material formed of *λίθος*, or *κράμια*, the occupation with *κράμια*.

Ibn Khaldūn, who lived in the 14th century, says that from the philosopher's stone a liquid or a powder might be prepared, called *iksir*, which, when thrown on molten copper converted it into silver, and molten silver into gold. In opposition to its etymology the word is here used for a liquid, because at that time *kimiyā* no longer meant the transmuting substance, but the science of transmutation, and explains why to-day we may understand by elixir a liquid.

We also find that the philosopher's stone is often called the red tincture, from *tinguo*, to moisten.

It appears, therefore, very probable that the name of our science is derived from *χυμός*, and the proper spelling would therefore be *Chymistry*, as the *Times* newspaper for a long time insisted upon. As, however, this derivation has not yet been proved quite satisfactorily, the time-honoured term *Chemistry* will remain in use, and I think be retained even if it should be shown that *χυμεία* was the original spelling.

Ordinary Meeting, December 2, 1879.

Dr. R. ANGUS SMITH, F.R.S., &c., Vice-President, in the Chair.

"On a Peculiar Feature in the Water of the Well in Carisbrooke Castle, Isle of Wight," by HARRY GRIMSHAW, F.C.S. The sample of the above water was taken by me on April 19, 1878, but was never opened or interfered with in any way until the following September. The water when taken was very bright and clear, and free from sediment of any description. It was totally devoid of odour, and was of a fresh and sparkling taste. The local features of the well from which the water derives its origin are as follows:—Carisbrooke Castle stands on a small isolated chalk hill, 239 feet above the level of the sea. The well is under cover in the "well house," and according to Jenkinson is 240 feet in depth. It is perfectly free, even at the surface of the water, from carbonic acid gas, or in fact of more than traces of any other gas than atmospheric air, as a candle floating on the water burns freely.

The bottle containing the water was opened on the 12th of September, and on doing so a very strong odour of sulphuretted hydrogen was perceived, and on testing with lead-paper an equally strong reaction for that gas was obtained. There was also apparent a slight sediment of a white colour not originally seen in the water. The reaction to litmus was perfectly neutral. The analytical data obtained, which were all that was possible from the quantity of water at command, were as follows:—

* Petermann, *Geog. Mitth.*, viii., 294.

† Kopp, *loc. cit.*, 209.

‡ Zosimus calls the substance by which copper is tinged yellow or converted into brass, *το δια της γαυρίας ἐξηριον*, a powder prepared by means of tutia; now tutia (zinc oxide) is still to-day used in medicine as a desiccative.

* Kopp, *loc. cit.*, 4.

† Kopp, *loc. cit.*, 76.

‡ *Zeitsch. Deutsch. Morgenl. Ges.*, xxx., 634.

Total solid matter ..	42'00	grs. per gal. (consisting of)
Mineral matter ..	22'40	" " and
Volatile matter ..	19'60	" "
Total hardness ..	12'30	" "
Magnesia hardness ..	1'70	" "
Chlorine ..	4'50	" "

The residue on heating blackened very much, and emitted a very strong unpleasant odour like burning animal matter.

The peculiarity of this water is of course the production of sulphuretted hydrogen, on standing for some time (in this case for five months) out of contact with the atmosphere. On leaving a small portion of the water in the bottle again corked up for some time the presence of sulphuretted hydrogen was not exhibited. This production of sulphuretted hydrogen proceeds undoubtedly from the reduction of the sulphates contained in the water by the excess of organic matter, and it is not unique in this instance, although it is not a fact of very common occurrence. I regret very much not having been able to bring back a sample large enough to admit of a determination of the albuminoid ammonia, and the nitrates and nitrites, as the quantity of these substances in a water of such a description would have been a very interesting item in the case, as touching on a point of great importance in pronouncing an opinion on the quality of a water. It is very possible and even probable that some chemists, given to judging for and against a water chiefly by indications of a single description, to the comparative neglect of many other analytical results, say by the amount of albuminoid ammonia, might actually fail to condemn a water such as the above; for many chemists appear to consider the ignition of the solid residue a comparatively unnecessary detail; and supposing even that the nitrates had been determined in this water, there are eminent chemists who have considered the presence of nitrates in deep well waters, in the chalk especially, as comparatively innocuous. The Carisbrooke well, as I have said, is in the chalky strata, and is 240 feet deep, and whether it contains nitrates or not, is, in my opinion, on the results above detailed, a most unfit water for potable purposes, and my reason for bringing the analysis of this water before the Society was to draw attention to the tendency often exhibited to draw the chief inferences of the quality of drinking waters from what may be called isolated reactions, without obtaining, or at all events without giving weight to, other indications, chemical and physical, which they exhibit.

Captain ABNEY, R.E., F.R.S., exhibited his photographs of the ultra-red portions of the solar spectrum, and first of all showed that the light transmitted by ordinary bromide of silver was of an orange tint, showing absorption in the lowest end of the spectrum. He then went on to explain how he had tried to load the molecules comprising this bromide of silver by means of gum resins, and that he had thus been enabled to photograph slightly beyond the lowest limit of the visible spectrum. Further researches proved that bromide of silver could be prepared in two molecular states, one that already shown, and the other in which absorption takes place in the red as well as in the blue. This was found sensitive to every radiation. He pointed out that the blue form of the silver bromide could be converted into the red form by simple friction, and that after friction it was insensitive to the ultra-red radiation.

Prof. ROSCOE here exhibited the different preparations of gold in minute division made by Faraday himself, some of which transmitted blue light and others red, showing that at all events two cases of molecular condition exist in the case of metallic gold.

Captain ABNEY then threw upon the screen photographs of the prismatic spectrum, in one of which the lowest limit of the prismatic spectrum was reached. He demonstrated this on the black board by setting up as ordinates the wave-lengths of the various portions of the photographs as obtained from the photographs of the diffraction spectrum.

He then exhibited various photographs of the ultra-red portion of the diffraction spectrum, extending from 7600 to about 11,000. He stated that the photographs from which he was making his final map were taken on double the scale, with twice the amount of dispersion.

He then showed various prismatic spectra, exhibiting different states of atmospheric absorption, in one of which Piazzi Smyth's rain-band was markedly visible.

After a short discussion, Captain Abney exhibited some photographs of the spectrum in natural colours.

NOTICES OF BOOKS.

Auxiliary Tables for Instruction in Chemical Analysis. (Hülf's Tabellen für den Chemisch-analytischen Unterricht.) Edited by OTTO WALLACH, with the co-operation of AUG. KÉKULÉ, A. BERNSTEIN, H. KLINGER, and C. WACHENDORFF. Bonn: E. Weber.

THE compiler of this little work fully recognises as not unfounded the prepossessions now widely entertained against the tabular form of instruction. He aims, however, at placing in the hands of the student not—as is too generally the case in tables—a mere routine to be followed, but a key to the understanding and intelligent execution of analytical methods. From the very outset the pupil is led to a consciousness of the reasons for selecting any particular procedure in preference to others. These objects are, we think, as far as possible within a space so limited, satisfactorily attained. The author recommends a close comparative examination of his Tables I. and X. The former of these contains preliminary remarks on a right understanding of the analytical process and on the use of the tables. The latter, No. X., is devoted to an examination of the solution containing metals of the "ammonium sulphide" group in presence of phosphoric or oxalic acid. In both these tables we find attention called to points often slurred over. Thus, as regards the preliminary step of dissolving the unknown body placed in the hands of the operator, be it a single compound or a mixture, we have met elsewhere with the direction to treat with water, or hydrochloric acid, or nitric acid, or aqua regia. Herr Wallach, on the contrary, advises these solvents to be used in succession in this order as far as needful, and supplies indications in case the substance, wholly or in part, does not dissolve in any of them. He also speaks of the examination of the physical properties of the sample, and of its behaviour in the dry way, not as mere tentatives which may be omitted at pleasure, but as rightly preceding the investigation in the wet way. At the foot of the table we find the advice "above all things to perform none of the operations without having a clear understanding of the reason why." The table for the examination of substances in water and acids, and of those where the elements present are masked by organic matter, strikes us, as very valuable.

The work being intended for beginners only, the rarer elements are not included in its plan, and the author does not count on rendering the oral advice and the supervision of the teacher unnecessary.

In fine, we may safely say that numerous as are the manuals, handbooks, and sets of analytical tables now in existence, the work before us will be found useful in the laboratories of our colleges, and is not to be set aside as an "Ilias post Homerum." There is no reference to examinations and the means of passing them, the author's object being simply to train sound chemists capable of doing good work.

The Use of Borax for the Preservation of Meat.—E. de Cyon and G. Le Bon.—The former author recommends the use of borax, and the latter pronounces it injurious.—*Biedermann's Centralblatt.*

CORRESPONDENCE.

FLUID MEDIUM FOR IMMERSION LENSES.

To the Editor of the Chemical News.

SIR,—Referring to my former letter on the subject of suitable fluids for use with microscopic objectives on the homogenous immersion principle (CHEMICAL NEWS, vol. xl., p. 213), I beg to call attention to a note appended to the recently published paper read before the Royal Microscopical Society,* June 11, 1879, by Professor E. Abbe, of Jena.

"Microscopists who are connected with chemical matters are kindly requested to promote the homogeneous immersion method by looking out for such less known liquids as afford some hope of being useful for the purpose, and I shall be glad to investigate every sample which may be sent to me for trial. A few drops only is quite sufficient for exact measurement by means of the refractometer of the refractive and dispersive indices."—I am, &c.,

W. T. SUFFOLK.

Stettin Lodge, St. Faith's Road,
Lower Norwood, S.E.,
December 15, 1879.

ALUMINIUM PLATE AS A SUPPORT IN BLOWPIPE WORK.†

To the Editor of the Chemical News.

SIR,—Critics no doubt differ in estimating the amount of acknowledgment due to inventors, but while cordially concurring with all you say regarding the merits of M. Landauer's lately published little work on blowpipe analysis, I would ask you, as one of our most distinguished discoverers, if you think it altogether fair to insert detailed descriptions of a man's invention in *thirteen pages* of a small book, as M. Landauer's English editors have made him do with regard to the new reactions on aluminium plate; to refer—not in the text, but in a note—to the inventor as only having "recommended" the novelty; and, finally, without further mention of the inventor's name, to refer thus to the matter in the index—"Hutchings: Aluminium plate reactions"? M. Landauer writes to me on the subject thus:—"I have, however, not stuck strictly to this principle [of not naming authors], but made exceptions with regard to the most important improvements. Among those is your application of Al plate." Having no copy at hand I cannot speak to the wording of the citation, but most likely it will be in harmony with the source used, viz., Hutchings's article, which said:—"By far the best support is the above-named Al plate, first used by Major Ross, and described in his work, 'Pyrology.'" But Hutchings also said, either in the *Mining Journal* or *CHEMICAL NEWS*, that "Al plate raised this entire branch of blowpipe analysis to a higher standard."—I am, &c.,

W. A. ROSS.

* London, W., December 14, 1879.

P.S.—The use of gold, by the way, for detecting nickel is not a novelty, but mentioned in Plattner's 4th edition.

CHEMICAL NOTICES FROM FOREIGN SOURCES.

NOTE.—All degrees of temperature are Centigrade, unless otherwise expressed.

Biedermann's Central-blatt für Agrikultur-Chemie.
November, 1879.

Application of a Natural Product as Manure.—Prof. F. Ullick.—The substances concerned are basalt and

* *Journal Royal Microscopical Society*, December, 1879, p. 823.
† The title used by Mr. Hutchings in *CHEMICAL NEWS*, vol. xxxvi., p. 208, 217.

the tufas and wackés found in its vicinity. The analysis of such a wacké yielded potash 0.68 and phosphoric acid 0.62. The author recommends the same material for absorbing crude potassium salts for application on the land.

Manurial Experiments for Determining the Requirements of Arable Soils.—Baron Dael von Koeth.—The results of experiments carried on for thirteen years near Maintz, on a calcareous soil. Reverted phosphates gave better crops than soluble phosphate employed alone. Gypsum was not appreciably useful.

Influence of Various Manures on the Combustibility of Tobacco.—Gaetano Cantoni.—The Stassfurt salts, night-soil, and all manures rich in chlorine injure the burning-power of the leaf.

Studies on the Digestive Process in Sheep.—Dr. E. Wildt.

Researches on the Digestion of Food in Horses under the Influence of Increased Work.—Prof. E. v. Wolff, Prof. W. v. Funke, Dr. C. Kreuzhage, and Dr. O. Kellner.

Digestion of Potatoes and Flesh-Meat in Pigs.—Prof. E. v. Wolff and Prof. W. v. Funke.

Chemical Studies on the Activity of Bees.—Dr. E. Erlenmeyer and Dr. A. v. Planta-Reichenau.—The authors conclude that wax is chiefly formed by the bees from sugar. Sugar occurs in honey chiefly as glucose, though cane-sugar is also present. Both the pollen of flowers and the saliva of the bee are powerful inverting agents.

Growth of Vines from Seed.—Dr. A. Blankenhorn.—The author maintains that the constitution of European vines is exhausted, owing to their continued propagation by layers and cuttings. He recommends propagation by seed as a defence against the *Phylloxera*.

Determination of the Value of Bone-Black.—Dr. Reinecke and G. F. Meyer.—The decolourising power of bone-black is inversely as its specific gravity.

Determination of Value of Wines.—MM. Houdart and Petit.—The authors determine the specific gravity at 15°, the alcohol at the same temperature, and the weight of the dry extract per litre.

Determination of Glycerin in Wines.—Prof. C. Neubauer and Dr. E. Borgmann.—The authors, using Pasteur's method as modified by Reichardt, found in genuine wines 0.978 to 1.667 per cent of glycerin.

Detection of Adulterations in Butter.—The methods for the detection of spurious fats are arranged under three heads: (1) microscopic examination, (2) determination of the specific gravity, and (3) separation of the fatty acids.

Presence of Sulphuric Acid in Milk.—Dr. G. Musso and F. Schmidt.—The authors have invariably detected sulphuric acid in normal milk.

Les Mondes, Revue Hebdomadaire des Sciences.
November 27, 1879.

Mechanical Equivalent of Heat.—Prof. von Waltenhofen.—The results obtained agree in a satisfactory manner with Joule's equivalent.

Laws of the Expansion of Metals.—M. de Heen.—The author has shown some time ago that in the metals of one and the same natural group the product of the coefficient of expansion by the absolute temperature of fusion is a constant number. He has recently enquired how the coefficient of the expansion of water varies with the nature and the quality (quantity?) of the substances held in solution, and he finds that there is also a constant relation between the coefficient of expansion of organic liquids belonging to one and the same homologous series and the boiling-point. The product of one of these magnitudes by the other is constant.

Chemiker Zeitung.
No. 49, 1879.

Manufacture of Permanent White, and Utilisation of the By-products.—The commencement of a lengthy memoir.

Manufacture of Magenta by Couplier's Process.—Dr. O. Witt.—The only manufacturers employing this method are Couplier himself and the firm of Meister, Lucius, and Brünig.

New Apparatus for the Determination of Urea.—G. Buts.—Requires the accompanying figure.—*Rép. Pharm.*, vii., 503.

MISCELLANEOUS.

University of London.—The following are lists of the candidates who have passed the recent Second B.A. and B.Sc. Examinations for honours:—

(B.A. and B.Sc. conjointly.)

Mathematics—First Class: R. S. Heath, B.Sc. (Scholarship), Trinity College, Cambridge; A. E. Steinthal, B.A., Owens College and Trinity College, Cambridge.

Mental and Moral Science—First Class: D. S. MacColl, B.A. (Scholarship), University College; W. H. Findlay, B.A., Merton College, Oxford; J. A. Newbold, B.A., Owens College. Third Class: M. A. Power, B.A., Stonyhurst College; D. C. Ross, B.A., University College; T. F. Kerry, B.A., Regent's Park and University Colleges; B. B. Le Tall, B.A., Owens College; W. A. Todhunter, B.A., private study.

(B.Sc. only.)

Chemistry—Second Class: E. H. Rennie, M.A., Sydney, St. Mary's Hospital and private study.

Experimental Physics—First Class: S. L. Hart, (Scholarship) St. John's College, Cambridge; S. G. H. Barfield,* private study. Second Class: J. W. W. Waghorn, private study; E. Hopkinson, Emmanuel, Cambridge, and Owens Colleges.

Botany—First Class: G. B. Hoffmeister, Gonville and Caius College, Cambridge. Third Class: J. G. Ridsdale, Guy's Hospital and private study.

Zoology—First Class: W. D. Halliburton (Scholarship), University College. Second Class: S. J. Hickson, Downing College, Cambridge; W. Overend, University College and St. Bartholomew's Hospital. Third Class: J. J. Fletcher, B.A., Sydney, Royal School of Mines and University College.

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